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The Design and Operation of a Real-Time Polynuclear Aromatic Hydrocarbon (PAH) Monitor for the Analysis of Combustion Products Formed in the Incineration of Navy Colored Smoke Compositions

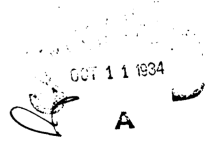
by Richard T. Loda Research Department

AUGUST 1984

NAVAL WEAPONS CENTER CHINA LAKE, CA 93555-6001



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FOREWORD

Polynuclear aromatic hydrocarbons (PAHs) can be produced during the thermal destruction of unserviceable colored smoke compositions. Because some of these compounds are carcinogenic, there is a need to monitor their possible release into the environment. The design, construction, and implementation of a real-time incinerator monitor system for PAHs is the subject of this report.

The work described in this report took place between December 1982 and October 1983. The work was performed with Pollution Abatement Research funds, Program Element No. 62765N, SEATASK Task Area Number WF65559 under the sponsorship of G. Young and under Task Area 50400 under the sponsorship of J. Short.

This work has been reviewed for technical accuracy by E. D. Erickson and C. E. Dinerman.

Approved by E. B. ROYCE, Head Research Fapartment 30 July 1984

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This publication describes design and construction of a fluorescencebased polynuclear aromatic hydrocarbon (PAH) monitor and calibration system. It also covers the installation and operation of this real-time monitor system during the incineration testing of Navy colored smoke compositions at the Los Alamos National Laboratory Controlled Air Incinerator facility, Alamos, N. Mex. During these tests, no PAHs were found to be present in the incinerator effluent gases at a concentration level 21 ppm (the approximate gas-phase detection limit of the monitor). Recommendations are listed for future system improvements. Comments assigned

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DISCLAIMER

The mention of specific equipment manufacturers, brand names, or specific contractors in this report does not imply their endorsement. Other commercially available equipment, having similar operational capabilities, or other qualified contractors, could be substituted for the equipment or contractors mentioned. Statements made in this document do not necessarily represent official Navy or DOD policies and practices; they are the best judgment of technical personnel, based on an experimental investigation of the problem, and a review of pertinent information. No statements in this document are intended to modify or replace any official safety SOPs, instructions, etc. Any prices listed were as of August 1983 and may not be the same now.

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INTRODUCTION

The Navy has an inventory of obsolete and unserviceable colored smoke compositions that must be disposed of in a safe, economical, and environmentally acceptable manner. Disposal by thermal destruction (incineration) has been demonstrated to be technically feasible (Reference 1), but this process, under certain conditions, produces polynuclear aromatic hydrocarbons (PAHs) from the incomplete oxidation of organic dyes present in the smoke compositions. A listing of the materials that have been found is given in Table 1. Since some of these PAHs are carcinogenic, effective control of PAH emissions must be achieved for the incineration process to be fully acceptable.

Pilot-plant scale experimentation is necessary to obtain a "best set" of incineration conditions to ensure the maximum destruction efficiency of these smokes, and a real-time monitor is required for the detection of PAHs in the combustion product. To accomplish these tasks, the Naval Weapons Support Center (NWSC), Crane, Ind., awarded a contract to the Los Alamos National Laboratory (LANL), Los Alamos, N. Mex., to experiment with incinerating the smoke compositions in their research incinerator. In addition, both the Naval Weapons Center (NWC) and the Battelle Columbus Laboratories (BCL), Columbus, Ohio (Reference 2) were contracted to design a monitor capable of detecting mixtures of PAHs on a one part-per-million (ppm) level. Ultimately, it was the PAH monitor system design of NWC that was implemented during the incineration tests undertaken at the LANL (References 3 and 4). If these tests prove successful, the Navy will subsequently build its own incinerator facility using the knowledge gained through these experiments.

This report describes the NWC's PAH monitor and calibration device design considerations. A listing of all of the company names and their addresses is included at the back of the report. It also covers the on-line installation and operation of this real-time system during the incineration tests performed at the LANL in September 1983. During these tests, no PAHs were found to be present in the incinerator effluent gases at a concentration level >1 ppm (the approximate detection limit of the monitor). Background material, specific to the PAH problem, will be discussed first. Following this, the experimental apparatus details will be given. This section covers the actual monitor design, before some of the results influencing it are presented.

TABLE 1. PAH Materials Found in the Combustion Product of Navy Colored Smoke Compositions.

PAH	Formula	Structure
Phenanthrene	C ₁₄ H ₁₀	
Anthracene	C ₁₄ H ₁₀	
Benzo(a)pvrene	с ₂₀ н ₁₂	∞\$
Benzo(e)pyrene	C ₂₀ H ₁₂	₩
Perylene	C ₂₀ H ₁₂	828
Chrysene	C ₁₈ H ₁₂	
Triphenylene	C ₁₈ H ₁₂	
Fluoranthene	C ₁₆ H ₁₀	∞ 8
Pyrene	C ₁₆ H ₁₀	
Naphthalene	c ₁₀ Hs	∞
1-Methyl Naphthalene	C ₁₁ H ₁₀	⇔

a Reference 1.

If this bothers the reader, it is suggested that the report be reviewed in the following order: the Introduction section (p. 3) through the Laser Inducted Fluorescence (LIF) System (p. 14); Results and Discussion section (p. 24) through the Comparison of PAH Spectra Static Cell section (p. 36); the LANL Incinerator Complex section (p. 14) through the PAH Monitor Design section (p. 24); and then the LANL Incinerator Tests section to the end (p. 36 through 75).

The results will essentially be presented in chronological order. Work on the flow and calibration system, with known PAH materials, will be described first. This will be followed by the evaluation of some commercially available instrumentation, and the choice of the monitor. Next, the PAH monitor system installation, and the on-line, real-time analysis will be discussed. Lastly, a series of recommendations, to improve the monitor operation for future work, will be presented.

BACKGROUND

Although PAHs are naturally present in many forms of vegetation and fossil fuel, the principal sources of ambient atmospheric PAH concentrations result from the incomplete combustion of hydrocarbons. Processes such as the combustion of coal, oil, gasoline, and diesel fuel, along with the operation of refuse burning power plants and wood burning stoves, all contribute significantly to the PAH pollutant problem.

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Since many compounds of the PAH class have been shown to be mutagenic and/or carcinogenic in animals, it is no surprise that a large body of work has been devoted to the study of these materials (References 5 and 6). Most analytical methods for PAH determination have been based on the isolation of individual PAHs in collected environmental samples through a variety of chromatographic techniques. Quantitative information is then gathered after the individual components have been identified. Quite often this entire process is carried out using a combination of instruments. For example, the gas chromatography/mass spectrometry (GC/MS) technique combines the high separation efficiency of gas chromatography with the high sensitivity and identification capabilities of mass spectrometry. High performance liquid chromatography with fluorescence detection is another popular instrumental combination for PAH analysis, and many other extensions of this basic idea are possible. Unfortunately, most of these procedures require time-consuming and troublesome collection, extraction, and concentration techniques. Sample recovery and the maintenance of sample integrity during the work up prior to analysis represent serious problems, and the desired information cannot be obtained in real-time.

The inherently high sensitivity of fluorescence detection, coupled with the intense fluorescence of many PAH compounds (References 7 through 11) makes fluorescence spectroscopy a powerfu' tool for the detection and identification of PAHs. Fluorescence spectroscopy has been used to measure PAH concentrations below 1 ppm, and this technique has the potential for making real-time measurements.

Fluorescence is a process in which radiation is emitted by molecules that have been previously excited by the absorption of radiation. When a sample is excited at a fixed wavelength $\lambda_{\rm ex}$, a fluorescence emission spectrum is produced by recording the emission intensity as a function of the emission wavelength $\lambda_{\rm em}$. Alternatively, a fluorescence excitation spectrum may be obtained by scanning $\lambda_{\rm ex}$, while recording the emission intensity at a fixed $\lambda_{\rm em}$.* These excitation and emission spectra are characteristic of the materials present in the sample. In the study of multicomponent mixtures, the ability of the analyst to select two wavelengths (excitation and emission) for the measurement of the fluorescing species leads to an enhanced selectivity for fluorescence detection over other techniques such as absorption spectrophotometry (since not all absorbing molecules fluoresce). This is especially true for the highly-fluorescing PAHs.

The problems associated with the separation and purification steps mentioned above have lead to an active effort to develop improved instrumental and mathematical curve fitting techniques for the cotal analysis of PAH mixtures without first isolating the individual components. The major problem area has not been sensitivity, but selec-The overlap of spectra from the different sample components and the inherently featureless nature of the individual spectra, especially when measured under ambient or higher temperature conditions, are the essential causes for this limitation. Nevertheless, low temperature site-selection spectroscopy (References 12 through 16), matrix isolation spectroscopy (References 12, 17 and 18), polarization measurements (Reference 19), time-resolved spectroscopy (References 20 and 21), selective excitation (Reference 22), derivative spectroscopy (References 23 through 28), selective modulation (Reference 23), synchronous excitation (Reference 29) and video fluorimetry (References 30 and 31) have all been employed to improve multicomponent analysis. A variety of numerical algorithms have also been used to deconvolute the overlapping spectra (References 32 and 33).

A fluorescence excitation spectrum is essentially an absorption spectrum, but measured indirectly by monitoring sample emission. The key point is that the sample cannot emit unless it first absorbs.

When we consider the specific requirements for a real-time PAH mixture analysis of stack gases, a number of the approaches mentioned above have definite drawbacks. Site selection and matrix isolation would require sample collection and cryogenic sample preparation. addition, these techniques, along with time-resolved spectroscopy, need relatively expensive laser sources for sample excitation. Cryogenic equipment and sample preparation are also required for polarization measurements, and selective modulation necessitates extensive modifications to standard spectrometers and the construction of additional electronics. In both selective modulation and synchronous excitation, the selectivity is enhanced, relative to simpler methods such as selective excitation, but the sensitivity is reduced because the data results from a convolution of the excitation and emission spectra. Derivative spectroscopy can enhance selectivity over selective excitation and is rather simple to implement, but the signal to noise (S/N) ratio is reduced by a factor of about 2 for each derivative taken This reduces the potential sensitivity of this (Reference 23). approach. Video fluorimetry can provide an entire excitation-emission matrix on a complex sample, in real-time, and is probably the most powerful technique of all those mentioned above. Its major drawback is that it requires the modification of a standard spectrometer and the purchase of an expensive (approximately \$50K), image intensified array detector, along with a computer controlled data acquisition system. Furthermore, even after one obtains and stores the vast amount of information contained in one, or multiple excitation-emission matrices, the subsequent analysis is by no means trivial. Sophisticated deconvolution techniques have been developed (References 32 and 33); but for complex samples, one still must know the total number of components in the mixture. This would not be easy to determine, a priori, for our particular problem. Therefore, one must question whether the additional instrumental cost would be justified in this case.

A real-time aromatic vapor monitor has been previously reported (References 26 and 28). It uses derivative ultraviolet (UV)-absorption spectrophotometry as its detection method and has been primarily successful in analyzing mixtures of smaller, highly-volatile aromatic compounds. Since it was expected that larger, less-volatile PAH compounds would be encountered in the smoke composition incineration process (Reference 1), and that interfering, non-fluorescent but absorbing species would also be present, it was felt that an absorption-based PAH monitor would not meet the program requirements.

As will be presented in greater detail, our PAH monitor design, which is based on the purchase and minor modification of a commercial fluorimeter, allows the capability to exploit absorption, selective excitation and emission, derivative spectroscopy, and synchronous excitation techniques. Complete excitation-emission matrices can also be acquired; not on the same time scale as video fluorimetry, but at a greatly reduced cost.

There is one final technique that has been used to detect PAHs that also deserves discussion. It is a simple spot test (References 34 and 35) that allows a person to visually detect the presence of <10 picograms of PAH in a collected sample. The sensitivity of this technique is a consequence of a process called sensitized fluorescence. Sensitized fluorescence can be thought of as an energy transfer between two materials. A donor molecule may be excited with UV radiation and transfer its excitation energy to an acceptor molecule whose excited state is lower in energy than the excited state of the donor molecule. This acceptor molecule can then fluoresce after having been indirectly excited (sensitized). The transfer of energy is most efficient when the acceptor molecule is present in a much lower concentration than the donor molecule. When a PAH is used as the acceptor in an appropriate mixture (PAH plus sensitizer), the limit of detection can be significantly lower than the non-sensitized fluorescence detection limit of the acceptor molecule (PAH) alone.

To perform this spot test on a sample extract, one makes three application spots on a filter paper with a microliter pipette. Two of the spots contain portions of the sample extract. A naphthalene sensitizing solution is applied to the third spot and one of the sample spots. After the spots have dried, they are observed under black-light (254 nm) illumination. Visual differences in intensity and color between the sample/sensitizer spot and either spot alone indicates sensitized fluorescence and the possible presence of PAHs. This spot test is very useful as a prescreen and can eliminate the need for costly analysis when no PAHs are present. Although this test cannot be used in a real-time fashion, and can only be used to detect the presence of PAHs as a class, its description has been included here because it is simple, rapid, sensitive, and quite inexpensive.

EXPERIMENTAL

MATERIALS

The PAH compounds that were used in the laboratory scale experiments and for calibration purposes during the LANL incineration tests were purchased from the Matheson Chemical Company and Eastman Chemical Company. They were used without further purification if their melting points and melting point ranges agreed with the quoted literature values. When discrepancies were found, the PAH compound in question was recrystallized from ethanol and the melting point remeasured until satisfactory results were obtained. The nitrogen gas used in the laboratory flow system experiments was generated from the boil-off of a liquid nitrogen source. The air used for the same experiments was filtered laboratory compressed air. No fluorescence background signal was seen from the air alone.

LABORATORY FLOW SYSTEM

The pyrex diffusion cell used to produce known gas phase concentrations of PAH material is shown in Figure 1. Its design is quite simple (Reference 36), and it has been used previously for the delivery of constant quantities of PAH vapor into a nitrogen gas stream (Reference 37). The solid PAH sample is introduced through a removable 1/4-inch outer diameter (OD) sample tube, which is connected to the main body of the diffusion cell with a brass Swagelok union. A Teflon ferrule is used to make the seal on the pyrex sample tube. The main body of the diffusion cell is fashioned from a 3.0 cm OD pyrex tube. The inlet and outlet ports are approximately 1 cm OD, and a 2.5 cm length of 0.1 cm inside diameter (ID) capillary tubing is used on the inlet side of the cell. The 3.0 cm diameter portion of the cell is 9 cm long, and the sample tube length is 11.5 cm. We took the average diameter of the sample tube to be 0.40 cm, including the Swagelok union. The sample tube length and diameter will be used later in the diffusion rate calculations as the average length of the diffusion path (1 = 11.5 cm) and the cross sectional area of the diffusion tube $(A = 0.126 \text{ cm}^2).$

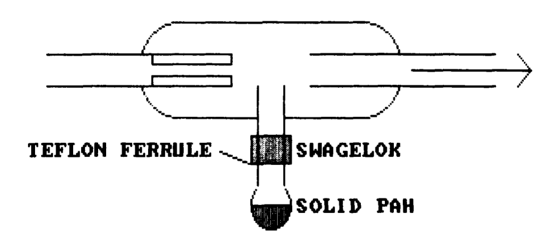


FIGURE 1. Diffusion Cell.

The diffusion cell is incorporated into the laboratory flow system as diagrammed in Figure 2. A 13-foot length of 1/4-inch OD copper tubing precedes the diffusion cell to allow the gases to achieve thermal equilibrium at the furnace temperature before picking up the PAH vapor being generated in the diffusion cell. The connection of the copper tubing to the diffusion cell is made via flexible stainless steel tubing, a Swagelok union, and a glass-to-metal transition tube on the diffusion cell. The purpose of the flexible tubing is to reduce the strain on the fragile glass components.

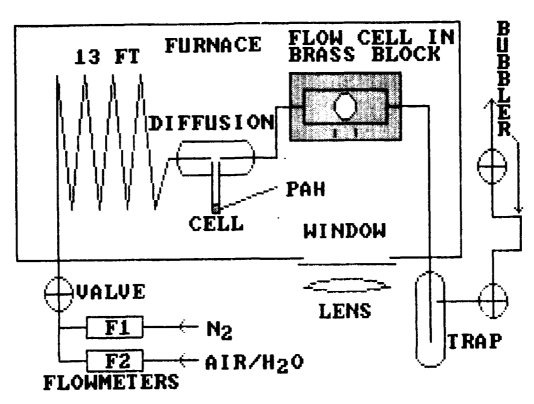


FIGURE 2. Laboratory Flow System.

After exiting the diffusion cell, the diluted PAH vapor stream then passes through a Suprasil-1 quartz fluorescence flow cell (1 x 1 x 2 cm), mounted within a brass block. Holes were made in the block so that the gas phase sample could be excited with laser radiation and the fluorescence emission could be observed at right angles to the excitation beam. A (f7 efficiency) lens collects the emitted light after it passes through a 2-inch diameter window in the insulated faceplate of the furnace. Quartz optics are used throughout to ensure high UV transmission.

The gas stream next flows out of the furnace through an ice/water trap which captures the PAH material. The connection from the fluorescence flow cell to the swagelok feed-through in the furnace faceplate is again made with flexible stainless steel tubing to reduce any strain on the fragile quartz flow cell. Finally, the gases pass through a mineral oil bubbler on the way to a chemical fume hood exhaust.

In order to set the gas flow rates and create gas mixtures, two Brooks Instruments, R-2-15-AAA flowmeters are connected in parallel. These flowmeters have metering valves at their inlet connections which are not shown in Figure 2. Before use, the calibration curves for these flowmeters were checked for accuracy with a bubble flowmeter.

The flow system was enclosed in a Forma Scientific Vacuum furnace (Cat. No. 3053). The door to the furnace was removed and replaced with an insulated faceplate containing the quartz window assembly. The vacuum gauge and valves were also removed, leaving a 1/2-inch diameter hole in the top of the furnace to be used as an access port for the laser beam. A quartz window was mounted over the hole to maintain thermal stability in the furnace. The maximum attainable temperature of the entire assembly is limited by the 232°C temperature rating of the Teflon ferrules. Higher temperatures could be achieved if these were replaced with a more heat resistant material, such as graphite.

Three chromel-alumel thermocouples are positioned in the furnace to monitor the temperature. The first is placed at the PAH sample tube, and the second on the main body of the diffusion cell. The third is placed inside the brass block. The brass block is also wrapped with a heating tape to allow some additional variac adjustment of the fluorescence flow cell temperature. The thermocouple potentials are monitored on three channels of a Hewlett-Packard model 3480A digital voltmeter, equipped with a model 3485A scanning unit, and converted to temperature readings using standard tables.

Before assembling the flow system, all the tubing was thoroughly flushed with several washings of methylene chloride. It has been our experience (Reference 38) that new copper tubing contains a significant number of PAHs, probably formed from the lubricants used during the manufacturing process. After assembly, but before use, the entire flow system was leak checked with helium using a Gow-Mac Instruments model 21-250 gas leak detector. This was done, not only to ensure system integrity, but also as an important safety check.

LASER INDUCED FLUORESCENCE (LIF) SYSTEM

The LIF system used for the laboratory scale experiments is diagramed in Figure 3. The flow system and furnace described previously are present in the upper right corner of the figure. The sample excitation is provided by a Lumonics TE 861T-3 rare gas halide excimer laser, operated on the XeCl line (308 nm). This laser can easily output 70 mJ in an 8 ns pulse at this wavelength, and it is also capable of generating a variety of other UV and vacuum-UV wavelengths when operated with other rare gas halide mixtures. The maximum pulse repetition frequency of the laser is 140 Hz.

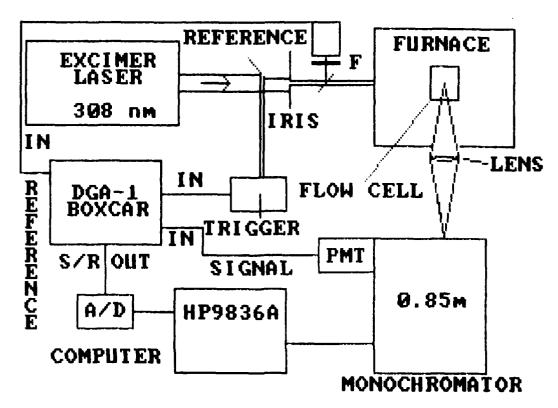


FIGURE 3. LIF System.

From the burn pattern of the excimer beam at 308 nm, its cross section is 4 x 20 mm. The excimer beam is irised to approximately 4 x 7 mm on its way to the furnace/flow cell and a small unused portion of it is reflected off of a pyrex plate to a photodiode trigger for the dual gated amplifier (boxcar). A quartz beamsplitter also picks off about 4% of the excimer beam and, after passing the beam through a 310 nm narrow bandpass filter and a Corning 7380 glass filter (F), sends the beam to a RCA 1P21 reference channel photomultiplier, The purpose of the reference channel is to correct the measured fluorescence emission signal for shot-to-shot laser intensity fluctuations and long term degradation of the laser power.

The beam is also reflected off of two mirrors which, for clarity, are not shown in Figure 3. The first is a 2-inch diameter CVI Corporation XC-2 dielectrically coated, >99% reflectivity, 308 nm mirror which is positioned before the beamsplitter and iris shown in the figure. The second is an ordinary aluminum mirror which is positioned on top of the furnace to steer the beam down through the access hole to excite

the sample in the fluorescence flow cell. Using the area factor for the iris, the maximum power to the sample would be on the order of 24 mJ/pulse. This estimate does not include beam divergence and reflectivity losses from the aluminum mirror and quartz plates. In the work described in this report, the actual power at the sample could have been as much as a factor of two lower than the value quoted above.

The emitted fluorescence is collected and focused into a Spex Instruments model 1404, 0.85 m, double monochromator equipped with 2400 g/mm holographic gratings and controlled by a CD2A Compudrive spectrometer driver. The dispersed emission then strikes the photocathode of an RCA C31034/76 photomultiplier inside a Products for Research model TE104RF thermoelectrically cooled photomultiplier housing mounted on the exit slit of the monochromator. The photomultiplier output is then sent to a Quanta Ray DGA-1 dual gated amplifier, and its ratioed output analog to digitally (A/D) converted for processing with a Hewlett-Packard model HP9836A desktop computer system. The CD2A Compudrive spectrometer driver is connected to the HP9836A computer via a RS-232 interface so that the whole experiment can be controlled from the HP9836A.

The data acquisition is handled under the software control of a program titled AD RLTIME (A/D real-time) written for these laboratory flow system experiments. The operator manually sets the monochromator to the emission wavelength to be monitored and enters information describing the PAH sample and the experimental conditions. The program then pauses, after displaying and labeling the axes for a fluorescence intensity versus real-time plot on the computer monitor screen. When all is ready, the program is continued by the operator and the data acquisition commences at time zero. The program plots the data and simultaneously displays the numeric values of the dependent and independent variables. Since each data point is the average of N samples over an operator prescribed sampling period, the standard deviation is also calculated and displayed, along with the point number, average intensity, and average time over which the samples were taken. A real-time interrupt function key can be pressed during the processing to record the point at which gas flowrates and/or concentrations are changed while the experiment is in progress. At the end of the data acquisition period the operator can then print the information on the experimental conditions and move a cursor through the plotted data to examine and print any points of particular interest (i.e., points taken near flowrate or gas mixture adjustments). Finally, the data set can be stored on disk for future use.

Under steady-state conditions, this program can also be used to generate a plot of fluorescence intensity versus wavelength by scanning the monochromator at a constant rate. As a check, the operator can use the real-time interrupt key to record at what point the monochromator

passes specific wavelength settings during the scan.* The AD_RLTIME program is listed in Appendix A, along with examples of the program output, which will be discussed further in the Results and Discussion section.

LANL INCINERATOR COMPLEX

The Los Alamos Controlled Air Incinerator (CAI) used for the incineration testing of the Navy colored smoke compositions has been described in great detail elsewhere (Reference 39). For our purposes, the relationship of the PAH monitor sampling location to the rest of the system is of primary importance. A schematic of the CAI is given in Figures 4 and 5, and a block diagram is presented in Figure 6. The

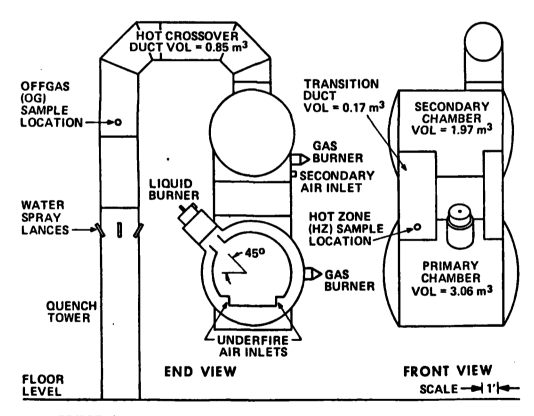


FIGURE 4. Schematic of LANL Controlled Air Incinerator.

We also have developed other, more elegant data acquisition programs that allow the HP9836A to have complete control of the monochromator scanning.

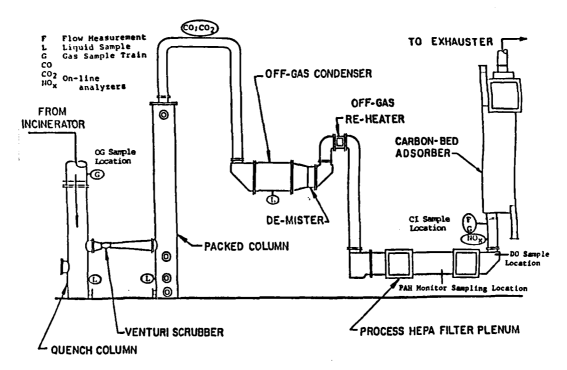


FIGURE 5. CAI Offgas System Sampling and Monitoring Locations.

HZ (hot zone, $T=982^{\circ}\text{C}$) sampling location is between the primary and secondary incineration chambers. The OG (offgas, $T=1093-1204^{\circ}\text{C}$) sampling location is just upstream of the spray quench column, and the DO (demister outlet, $T<93.3^{\circ}\text{C}$) sampling location is downstream of the offgas demister unit, just after the two High Efficiency Particulate Air (HEPA) filters. The PAH monitor sampling location ($T<93.3^{\circ}\text{C}$) is in between the HEPA filter assemblies, just before the DO sampling location.

Entropy Environmentalists, Inc., were contracted to perform offgas sampling at the HZ, OG, and DO locations during all phases of the incineration tests. They also recovered and prepared the samples for subsequent analysis by independent laboratories. For the details of the gas sampling equipment, procedures, reporting, and scheduling one should refer to their offgas sampling work plan (Reference 40).

The feed liquid for the incineration tests is a mixture of Navy smoke composition, fuel oil, water, and a wetting agent. It is injected into the incinerator after being dispersed with an atomization nozzle. Several different smoke compositions were fed into the CAI

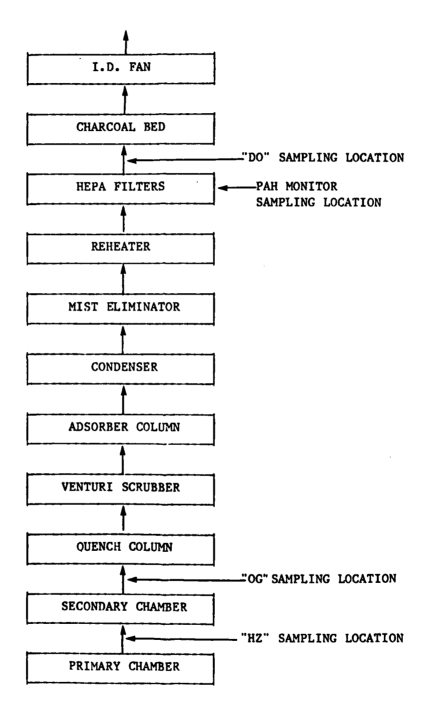


FIGURE 6. Incinerator Air Flow Schematic Showing Sampling Locations.

over 11 phases of operation. The feed schedules and operating conditions are all detailed in the Los Alamos test plan (Reference 41). Chemical compositions of the Navy colored smokes investigated have been reported elsewhere (Reference 1).

Comparative analysis of the real-time results obtained with the PAH monitor system and the collected samples will be made when data from the collected samples become available. At that time, the PAH monitor results will be compared with the data collected at the DO sampling location since these data sets were obtained under nearly equivalent sampling conditions.

PAH MONITOR DESIGN

Basic Spex Fluorimeter

The Spex Instruments, Inc., FLUOROLOG 2 series of fluorimeters are modular in their design. This allows one to choose among a variety of excitation sources, spectrometers, sample compartments, detectors, and accessories to assemble an instrument for a particular application. An optical schematic of a model F112 version of this instrument is presented in Figure 7. This figure is a composite drawing, pieces of which were taken from the Spex advertising literature (Reference 42). The F112 designation defines the spectrometer and sample compartment make-up of the instrument. As can be seen from the figure, there is a single (F1) excitation spectrometer which disperses the light emitting from a xenon lamp source. The spectrometer grating and slits determine the wavelength and bandpass of light used to excite the sample, which is located in the single-beam (F 1) sample module. The emitted fluorescence is then collected in either a right angle (90 degrees) or front face (22.5 degrees) geometry (user selectable), and dispersed in a double (F 2) emission spectrometer before striking the photocathode of a photomultiplier tube mounted on the exit slit of that spectrometer. If desired, other versions of the instrument (e.g., F122, F212, and F222, etc.,) can be configured or the existing system upgraded.

The Spex DATAMATE is a microprocessor (Motorola 6800) based spectrometer controller and data processor for the FLUOROLOG 2 series. A listing of its rather extensive capabilities is given in Appendix B. It is important to note that one can easily accomplish selective excitation and emission, synchronous excitation, derivative spectroscopy, and acquire complete excitation-emission matrices with the standard fluorimeter.* DATAMATE software programs can also be written for fully automatic operation of the instrument.

Refer to the Background section for the relevance of these methods to the problem of PAH analysis.

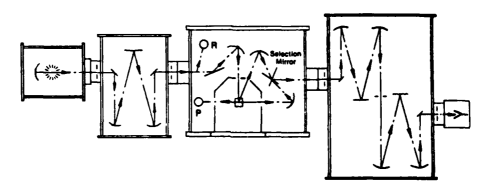


FIGURE 7. Optical Schematic for the Spex Industries, Inc. FLUOROLOG 2 Version F112.

The specific components which comprise the model F112 fluorimeter purchased for this project are given in Table 2. Comments regarding the individual components in relation to the overall PAH monitor design considerations are given below, in order of the item numbers in the table:

- 1. A single excitation monochromator was chosen, rather than a double, in order to maximize the throughput of the lamp. Since fluorescence intensity depends directly upon the excitation intensity, greater throughput of the lamp should translate to a greater sensitivity of the PAH monitor.
- 2. A double emission monochromator was chosen because of its superior stray light rejection as compared to a single monochromator. It was felt that scattered light might be a problem if particulates were encountered in the work.
- 3. We chose the cheaper fixed slit sets over the more expensive micrometer adjustable ones realizing that there would probably be little need for adjustments because of the broad nature of the PAH fluorescence.
- 4. The excitation grating was blazed at 250 nm to maximize the lamp throughput for wavelengths known to excite the PAHs of primary interest.
- 5. The two emission gratings were blazed at 300 nm to maximize the fluorescence throughput for those wavelengths over which the PAHs emit.

TABLE 2. FLUOROLOG 2 Model F112.

Item	Qty	ty Cat. no. Description						
1	1	1681B	Single excitation monochromator, 0.227 m f/4					
2	1	1680B	Double emission monochromator, 0.227 m f/4					
3	4	1679	Sets of fixed slits					
4	3	16016	Gratings, 1200 g/mm, in kinematic mounts. Excitation blaze = 250 nm, emission blaze = 300 nm					
5	1	1691	Single-beam sample module					
6	1	1909	Xenon lamp housing					
7	1	1907	Xenon lamp, 450 watt (W)					
8	1	1907P	Power supply for 1907					
9	1	1914G	Cooled R928 photomultiplier assembly					
10	1	1910	Quantum counter reference assembly					
11	1	DM1A	DATAMATE, with DM101 and DM104					
12	1	DM101	Input channel (additional)					
13	1	DM102	Photon counting (PC) acquisition module					
14	1	DM103	Direct current (DC) acquisition module					
15	1	DM104	High voltage (HV) power supply (additional)					
16	1	DM105	Data and programming package					
17	1	DM111	Dual disk drive					
18	1	DM112	Digital plotter					
19	1	1931A	Brass sample heater/cooler block					

The total cost, including installation and General Services Administration (GSA) discount, was \$36K.

^{7.} A 450-watt xenon lamp was chosen rather than the standard 150-watt lamp to again maximize the excitation (and therefore emission) intensity.

^{9.} The cooled photomultiplier housing was chosen in order to reduce the dark count (thermal noise) of the photomultiplier. The relatively inexpensive R928 photomultiplier was chosen because its spectral response, over the wavelength range germane to the PAH problem, is actually better than some more expensive tubes.

- 10. This assembly enables one to ratio the lamp intensity to the fluorescence emission intensity. Thus, one can correct the fluorescence excitation spectra for the fact that the lamp intensity is a function of wavelength. It also corrects for errors caused by lamp intensity fluctuations during a scan.
 - 12. Needed for 10.
- 13. PC is much more sensitive than DC measurement for weak fluorescent signals.
- 14. The DC module is for the reference channel. The more expensive PC module is not needed here because plenty of light gets to the reference channel (R in Figure 7).
 - 15. Needed for 10.
 - 17. Needed for program and data storage.
 - 18. Needed for hard copy output.
 - 19. Needed to house and heat the fluorescence flow cell.

Flow Cell Modifications

In order to perform the real-time fluorescence analysis of the incinerator effluents or calibration cell samples with this fluorimeter a minor modification of the standard F112 sample chamber is required. The fluorescence flow cell assembly constructed for this purpose is diagramed in Figure 8. An NSG Precision Cells type 501 FL UV flow cell (\$185.00) is mounted within the Spex 1931A brass sample heater/cooler The hose connectors on the 1931A were drilled out to allow for the insertion of two, 100-watt Chromalox cartridge heaters (CIR-1012). The heaters are controlled with a Chromalox model 3912 digital indicating on-off temperature controller, and the feedback is provided by a type J thermocouple (iron/constantan). Initially, with the thermocouple positioned in a hole at the top of the brass block, a range of 7.2°C was measured during the on-off cycle of the controller. By positioning the thermocouple in a hole in the brass block at the center of one of the cartridge heaters and wiring the heaters in series with a resistor, we were able to reduce the temperature fluctuations to an acceptable ±1.4°C at a set point of 65.6°C.

The connection from the flexible stainless steel tubing to the quartz flow cell was made with a short length of 1/4-inch OD Teflon tubing. This size fits nicely over the 5 mm OD flow cell ports and can be softened with a heat gun to stretch over the stainless steel tubing.

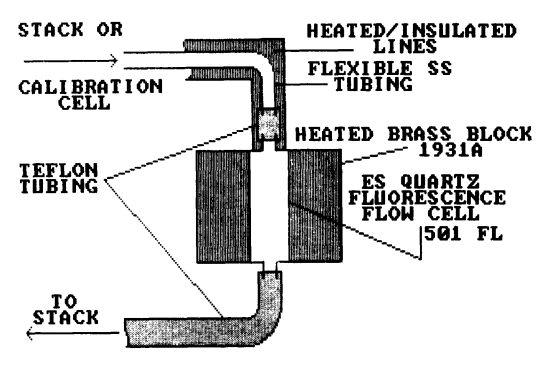


FIGURE 8. Fluorescence Flow Cell.

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The 232°C rating of the Teflon limits the maximum temperature of the assembly, but given the temperature range of the stack gases at the PAH monitor sampling location (<93.3°C), the Teflon range is more than adequate. The flexible Teflon tubing is also used on the exit side of the flow cell, and this part of the line is unheated. Variac controlled heating tapes maintain the temperature of the incoming line, which is also wrapped with insulating tape. The temperature of the line is monitored with a type J thermocouple (see next section for more details).

The incoming and exiting lines, along with the cartridge heater wires and thermocouple, all fit through holes which are already in the Spex 1691 sample module. The 1931A brass block is made to mount into the sample module such that the flow cell is optically aligned with the rest of the instrument. No additional adjustments are necessary. This means that the flow system can be very easily dismantled at the Teflon joints for cleaning, repair, etc. Also, since the brass block will hold standard 1 cm cuvettes, the system can be easily reconfigured for the fluorescence analysis of liquid (or solid) samples. To do this, one need only cool down the block and flow lines, stop the flow, dismantle and remove the flow cell, and replace it with the sample of interest contained in a standard 1 cm cuvette. This could be done to analyze for PAHs in collected and extracted samples from other parts of the incinerator system.

PAH Monitor Flow System

The overall PAH monitor flow system incorporating the fluorescence flow cell assembly of Figure 8 is given in Figure 9. The stack gases from the PAH monitor sampling location (refer to Figures 5 and 6) pass through 11 feet of heated and insulated tubing (lines 1 and 2) before entering the fluorescence flow cell, mounted in the Spex instrument. A calibration cell (line 3) is teed into line 2 so that the instrument response to a known PAH concentration can also be measured. The length of lines 3 and 2, from the exit of the furnace to the entrance of the fluorescence flow cell, is 8 feet. All the heated and insulated lines are made from 1/4-inch diameter stainless steel tubing which was cleaned with acetone before use. Variac controlled heating tapes maintain the temperatures in the heated lines, and type J thermocouples are positioned at the points labeled 1-4 in the figure. An Omega Engineering model DSS-650 type J thermocouple thermometer monitors the temperatures at those locations.

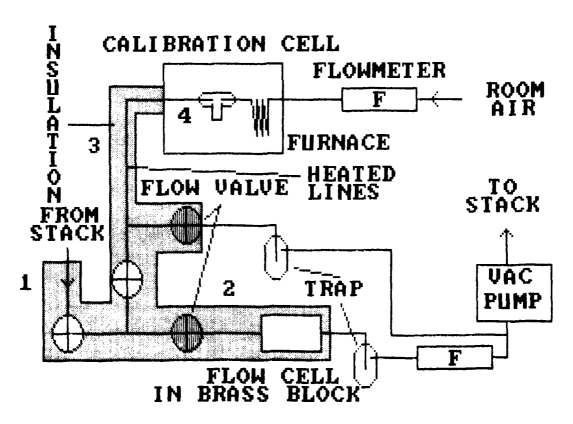


FIGURE 9. PAH Monitor Flow System.

The line running parallel to line 2 (and teed into it just before the vacuum pump) permits the calibration cell to continue functioning normally when the incinerator gases are being analyzed. If it were not there, PAH materials in the heated furnace would continue to diffuse and eventually plate-out on line 3 and the line going to the calibration cell flowmeter. This would defeat the purpose of the calibration cell and could expose workers to PAH vapors coming from the flowmeter inlet.

The unheated lines are made of either Teflon or polyethylene tubing, and an ice-water mixture is used to cool the traps. The flow-meters are Gilmont Instruments Cat. No. F1100 size #1 (maximum flowrate 270 ml/min, standard air at one atmosphere and 70°F). They are connected in series (for line 2) so that one can check for leaks in the monitor system by observing the relative flowrates on the two flow-meters. A modest vacuum pump is needed to generate the flow because the Los Alamos CAI operates at a negative pressure (50 inches of water less than atmospheric at the site). At negative pressures, any leaks in the incinerator system would cause materials to leak in, not out. This is done as a safety precaution, which also permits the processing of radioactive waste materials on other contracts.

To check the instrument response to a known PAH concentration, the system is operated with the on/off valve from the inciner tor stack in the off position. The on/off valve from the calibration call is in the on position, and the flow valve for the line parallel to line 2 is closed. In this configuration, ambient room air from the area near the furnace is drawn through the flowmeter under the action of the The flowrate is set by the flow valve just before the The 10-foot coil of tubing inside the fluorescence flow cell. Fisher model 556 laboratory furnace allows the air to equilibrate at the furnace temperature before entering the diffusion cell (refer to Figure 1).* The air stream dilutes the PAH vapor being generated in the diffusion cell, and the mixture flows through the heated lines to the fluorescence flow cell for analysis. After exiting the flow cell, the PAH material is collected in the trap, and the air stream passes through the second flowmeter. Following its passage through the pump, the air is returned to the incinerator stack well beyond the DO sampling location (refer to Figure 5).

To analyze the incinerator stack gases, one simply opens the on/off valve from the stack, closes the on/off valve from the calibration cell, and opens the flow valve in the line parallel to line 2. This

An Ultratorr fitting with a Viton O-ring was used at the LANL, rather than a Swagelok fitting with a Teflon ferrule, as shown in Figure 1.

keeps the calibration cell functioning while the stack gas analysis proceeds. The flowrate in the parallel line need only be enough to keep the PAH material in the calibration cell from plating out in the lines. The flowrates in both lines are set by their respective flow valves. If, at any time, the operator wishes to recheck the calibration, the valves are simply reconfigured as described in the last paragraph.

RESULTS AND DISCUSSION

DIFFUSION CELL EVALUATION

Theory

The diffusion coefficient for two perfect gases, designated 1 and 2, can be calculated using the following expression (References 43 and 45):

$$D = \frac{3}{8(2\pi)^{1/2}N} \times \frac{(RT)^{3/2}}{P} \times \frac{\left(\frac{M_1 + M_2}{M_1M_2}\right)^{1/2}}{d^2}$$
(1)

where

D = diffusion coefficient in cm²/(sec-molecule)

 $N_{\rm O}$ = Avagadro's number = 6.023 x 10^{23} molecules/mole

 $R = ideal gas constant = 8.31448 \times 10^7 (dyne-cm)/(mole-oK)$

 $T = temperature in {}^{\circ}K$

P = total pressure in dynes/cm² (1.01325 x 10⁶ at 760 torr)

 M_1 = molecular weight 1 in g/mole. (N_2 , air, etc.)

 M_2 = molecular weight 2 in g/mole. (PAH)

 $d_{12} = 1/2(d_1 + d_2)$ in cm

 d_1 = molecular hard sphere diameter for vapor 1 in cm

d₂ = molecular hard sphere diameter for vapor 2 in cm

One can make an estimate of the hard sphere collision diameter from data on the bond lengths and angles of the molecules involved. Using values of 1.4 and 1.1 Å for the aromatic C=C and C-H bond lengths, 1.1 Å for the N_2 bond length, and the cosine of 30 degrees, $d_{12} = 5.14$ Å can be easily computed for the anthracene- N_2 combination. A geometry where the long axes of the molecules were placed end to end was assumed in the calculation.

The vapor concentration delivered by the diffusion cell is determined by the ratio of the rate of diffusion of vapor from the diffusion tube and the diluent gas flow rate. The diffusion rate can either be measured from the change in weight of the sample tube, or calculated from the diffusion equation (Reference 36):

$$r = \frac{DAM_2P}{RT1} \times ln \left(\frac{P}{P-P}\right)$$
 (2)

where

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r = molecular diffusion rate in mg/(sec-molecule)

D = molecular diffusion coefficient in cm²/(sec-molecule) (from equation 1 above)

A = cross sectional area of diffusion tube in cm²

 M_2 = molecular weight of vaporizing species in g/mole

R = gas constant = 0.08205 (1-atm)/(mole-oK)

T = temperature in °K

1 = average length of diffusion path in cm

P = total gas pressure in diffusion cell in atm

p = partial pressure of vapor at temperature T

Equations 1 and 2 are incorporated in a BASIC program called DIFF RATE (Diffusion-Rate), listed in Appendix C along with sample output (which should be rounded to 3 significant figures). The program contains the parameters specific for the flow system experiments done at the NWC, i.e., P = 700 torr, l = 11.5 cm, and A = 0.126 cm². Two other versions of this program are parametrized for sea-level, and the LANL.

They are called DIFF_SEA and DIFF_LASL, respectively, and sample output is also included in Appendix C for comparison purposes. A program equivalent to DIFF_LASL, but written in the Spex Datamate programming language, is presented in Appendix D. All these programs permit the calculation of the PAH vapor concentration delivered by the diffusion cell under a variety of conditions.

PAH Diffusion Rate Measurements

In order to determine the accuracy of the simple calculations outlined previously, experimental measurements of PAH diffusion rates were undertaken, using the laboratory flow system of Figure 2. To do this, the cleaned diffusion cell sample tube was first loaded with approximately 10 mg of solid PAH. Next, this tube was weighed on a Mettler HL52 microbalance, and replaced in the flow system. At this point, the time was recorded and the furnace was heated to the desired operating temperature. A N2 diluent gas flow was also initiated to keep the diffusing PAH vapor from condensing in the flow lines (nominally 40 ml/min). The system was maintained at a steady state for such a time as to ensure a measurable weight loss in the sample tube. After the prescribed time period had elapsed, the furnace was turned off and the total time recorded. When room temperature was achieved, the No flow was stopped, and the sample tube reweighed. The mass loss, divided by the total time, gives a measure of the appropriate PAH diffusion rate for that set of experimental conditions. This value can then be compared to that calculated using equations 1 and 2.

To minimize the error caused by the finite time required to achieve steady state operating conditions, the start and stop times were defined as the times when power to the furnace was started and stopped. In this way, the "lack" of material diffused while the furnace was heating up was partially corrected for by the "extra" material diffused as the furnace was cooling down. Also, each experiment was run over a total time period of several days. Therefore, the approach of the diffusion cell to the steady state (a few hours) represented a small error.

The results presented in Table 3 show that the simple calculation can reasonably predict the measured diffusion rate for a number of PAHs over a range of conditions. In the case of the largest discrepancy between measured and calculated values (pyrene), it is quite likely that the biggest problem was in the accuracy of the pyrene

TABLE 3. PAH Diffusion Rate Measurement.

РАН	Anthracene	Phenanthrene	Naphthalene	Pyrene
T (°C)	159.0	133.5	25.0	128.0
p (torr) (Reference 46-48)	2.25	2.03	0.087	0.105
d ₁₂ (A)	5.14	5.15	3.9	5.15
M ₂ (g/mole)	178.0	178.0	128.0	202.0
Weight loss (mg)	7.73	5.73	0.20	0.51
Time x 10^{-5} (sec)	3.723	2.628	1.980	4.968
r x 10 ⁵ (mg/sec) measured	2.08	2.18	0.101	0.103
r x 10 ⁵ (mg/sec) calc. (DIFF_RATE)	2.28	1.98	0.0935	0.115

 $^{^{}a}$ N₂ diluent gas (M₁ = 28 g/mole) at P = 700 torr (NWC).

vapor pressure parameter. The value used had to be extrapolated* from literature data measured over a 69 to 85°C range (Reference 48). In addition, some laboratory temperature fluctuation could also have occurred during the naphthalene experiment, thereby affecting its accuracy. The largest difference between measured and calculated values encountered was approximately a factor of two, and this "worst case" measurement was purposefully made using impure phenanthrene.

LABORATORY FLOW SYSTEM EXPERIMENT USING LIF

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When we were ready to begin our flow system experiments, the research group at Battelle Laboratories had already measured the LIF emission spectra of a number of the PAHs listed in Table 1. These observations were made with the PAHs at elevated temperatures in a

The literature data was fit to an equation of the form $\log_{10}P = -a/T + b$ to determine values for a and b. With these values, P could then be calculated at another T. For accuracy, the T value, for which the new calculation of P is required, should be close to the measured data range.

static nitrogen atmosphere. The question of fluorescence quenching by oxygen and other atmospheric gas components (water, carbon dioxide, etc.) had not yet been addressed. Quenching would significantly reduce the measurable fluorescence intensity for a given PAH concentration, which in turn, would raise minimum detection limits of the fluorescence technique. Since this information was critical to the monitor design considerations, flow system experiments were conducted with the apparatus shown in Figures 2 and 3. The intent was to complement the LIF work already begun at Battelle.

The first molecular combination studied in the flow system was anthracene-N2. The RUN #1 output, given in Appendix A, shows the LIF emission spectrum of anthracene taken at 164°C (fluorescence cell This spectrum was taken under steady state conditions temperature). while scanning the wavelength drive of the monochromator at a constant Therefore, the time axis is also the wavelength axis. From the DIFF RATE sample calculation, given in Appendix C, the anthracene concentration was calculated to be 7.47 ppm at the 40 mL/min N2 flowrate. Using the data in Table 3, the calculated value should be adjusted by a factor of 2.08/2.28 = 0.912, which gives a measured value of 6.81 ppm. This LIF emission spectrum, obtained with a flow system, agrees quite well with the static cell measurement given in Figure 3 of the Battelle report (Reference 2), although their results were obtained at much higher concentrations (210 - 500 ppm). The excimer laser at 308 nm does not excite into the absorption maximum for anthracene at 344 nm. Using the excimer pumped dye laser at this wavelength could increase the LIF intensity by about a factor of 7. In addition, the broad emission spectrum does not require the 0.23 nm resolution used. throughput (and therefore sensitivity) could be increased by increasing the bandpass of the monochromator. The use of a smaller, higher throughput (and cheaper) monochromator could increase the sensitivity even further, if desired.

The output of RUN #2, given in Appendix A, shows the results of flow changes on the anthracene- N_2 LIF intensity at fixed excitation and emission wavelengths. The calculated anthracene concentration can be obtained from the DIFF RATE output in Appendix C. The intensity versus time plot shows that the intensity response is directly proportional to concentration, i.e., a change of a factor of two in flow rate (concentration) produces a change of a factor of two in the LIF intensity. Also, after each change in flow rate, the steady state is reached in a time period given approximately by $1^2/2D$ (Reference 36). These results demonstrate, quite nicely, how easily known concentrations of PAH material can be generated with the flow system/diffusion cell combination.

The effect of laboratory air on the anthracene LIF intensity is presented in RUN #3 of Appendix A (with appropriate calculations in

Appendix C). Air was found to quench the LIF intensity by about a factor of 5.5. The comments regarding the problems in setting the flow rate in the AD RLTIME output listing meant that one needed to keep an eye on the flowmeter and make minor adjustments with time, not that there were major deviations of the flow. When the change was made from N₂ to laboratory air at the same flow rate, the response time for the LIF intensity change reflected the time required for the new diluent gas to travel through the volume of tubing before reaching the fluorescence flow cell. Flow rate changes, on the other hand, were reflected in immediate intensity changes, as with the previous experiments. Our air quenching results were consistent with those on other PAHs studied by Battelle (Reference 2), and an order of magnitude was found to be a reasonable quenching estimate for the PAHs as a class. The largest value observed was for naphthalene (times 20, taken from Figures 21 and 22, Reference 2). This makes intuitive sense in that naphthalene has a relatively long fluorescence lifetime and would therefore undergo a larger number of collisions in the excited state before emitting a photon. This would increase the probability of its fluorescence being quenched relative to a molecule like anthracene, whose fluorescence lifetime is shorter. It was not felt that the magnitude of the air induced fluorescence quenching of the PAHs was large enough to warrant any basic PAH monitor design changes.

In examining the standard deviation data for all three runs, and comparing it to the graphic plots, the magnitude of the "noise" in the data was initially perplexing. The approximately 0.5 volt intensity fluctuations in the LIF, versus the <0.1 volt standard deviations (over the 20 data reads per second), were traced to the mineral oil bubbler in Figure 2. The bubbling action of the gas was causing pressure (and therefore concentration) fluctuations in the fluorescence cell. This phenomenon was not discovered early in the testing because no fluctuations were seen on the flowmeters, where one would expect to observe them. The 13 feet of tubing in the flow line acted as a ballast on the inlet side of the system, which damped them out. Also, the capillary tubing on the inlet side of the diffusion cell (Figure 1) would prevent the pressure backup to the flowmeters. Taking the data in the plots as an estimate, concentration fluctuations on the order of 500 parts-perbillion (ppb) were being observed in these experiments. The mineral oil was subsequently removed from the bubbler.

A program update meeting was held at LANL in April 1983. The reason for holding the meeting there was to familiarize all those concerned with the CAI site, where the incineration testing of the Navy colored smokes would be conducted. One of the outcomes of this meeting was the decision to move away from the laboratory LIF oriented experiments and towards the construction of a PAH monitor utilizing fluorescence detection. Our work shifted to the evaluation of commercially available equipment for this purpose. This topic will be discussed in the next section.

COMPARISON OF PAH SPECTRA (STATIC CELL)

Beckman DU-7 Absorption Spectrophotometer

In the process of developing a suitable test for the ability of commercial instrumentation to detect gas phase PAH concentrations, it was decided to measure the absorption spectrum of some possible candidate materials. One such spectrum of a mixture of naphthalene and phenanthrene is given in Figure 10. A small crystal of each material was carefully placed in the bottom of a cuvette (the special care was taken to make sure none of the solid material stuck to the cuvette windows). The cuvette was capped, placed inside a Beckman DU-7 absorption spectrophotometer, and the UV/VIS spectrum was obtained. The concentration of the gas phase material was determined by the partial pressure of each component. The spectrum of Figure 10 is essentially that of naphthalene alone since it is identical to a similar spectrum obtained with no phenanthrene in the cuvette. The instrument is incapable of detecting phenanthrene under these conditions.

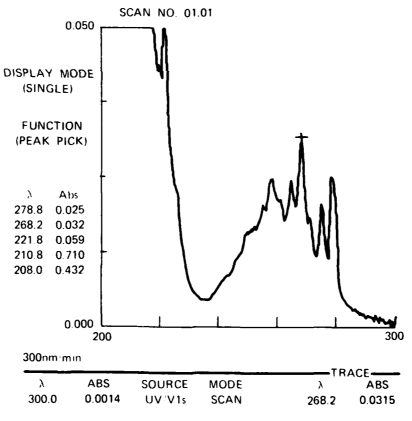


FIGURE 10. Absorption Spectrum of a Naphthalene/Phenanthrene Mixture, Taken in Air, at 23°C. Pathlength = 1.0 cm.

From the absorbance, pathlength, and the molar extinction coefficient, the naphthalene concentration can be calculated. Using an approximate value of 6×10^3 L/(mole-cm) for the extinction coefficient of naphthalene near 286 nm (Reference 49), a value of 5.25×10^{-6} molar can be calculated for the concentration, given the absorbance value from the figure. Multiplying by the molecular weight, and using the conversion factors in Appendix D, this translates into approximately 138 ppm. "Approximately" is used here because the extinction coefficient value was taken from a solution spectrum and because the background and absorption spectra were not taken in matched cells. Nonetheless, it is a reasonable estimate and an independent check on what follows below.

The concentration can also be calculated with the following expression:

$$ppm = (p/P) \times 10^6 \tag{3}$$

where

ppm = parts-per-million

p = partial pressure of vapor

P = total gas pressure

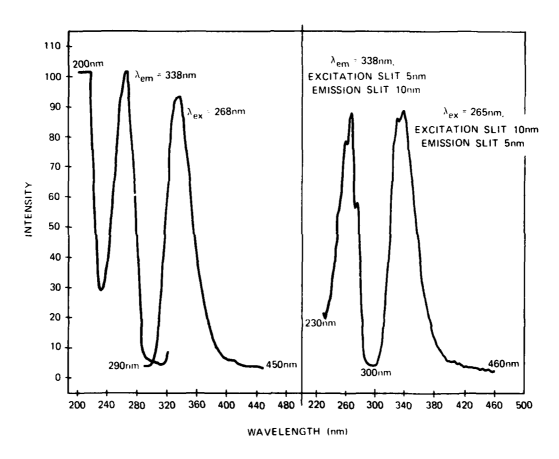
At 23°C, the vapor pressure of naphthalene is 0.0712 torr (Reference 46). At a total pressure of 700 torr (NWC), equation 3 gives a value of 102 ppm. Given the considerations mentioned above, there is reasonable agreement with the absorbance-derived result. For phenanthrene, with a vapor pressure of 1.39 x 10^{-4} torr (Reference 48), the concentration would be 0.2 ppm. It is not surprising that it went undetected.

In all fairness to the absorption technique, it should be mentioned that a long pathlength cell (e.g., 10 m) can be added to an absorption instrument, greatly increasing its sensitivity. Unfortunately, this still does not solve the selectivity problems discussed in the Backgound section. Also, alignment of a 10 m cell requires a certain amount of patience, and it can be difficult to clean, if contaminated.

Perkin-Elmer MFP-44B Spectrofluorimeter

Figure 11 shows the fluorescence excitation and emission spectra of the same naphthalene/phenanthrene sample studied above. Again, the 0.2 ppm phenanthrene component was not detected, but the signal to

noise ratios for the naphthalene spectra were somewhat better than those in Figure 10. The emission spectra compare quite favorably to the LIF results of Battelle (Reference 2), and we now have a selective excitation capability. Because this sample was studied in air, the fluorescence quenching factor is also included in the measurement. Extrapolation of these data would indicate that one could detect naphthalene down to about a 10 ppm level.



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FIGURE 11. Fluorescence Excitation and Emission Spectra of a Naphthalene/Phenanthrene Mixture, Taken in Air, at 23°C. Pathlength = 1.0 cm.

As a further test of the instrument, a cuvette containing phenanthrene was mounted in a brass block which fits inside the Perkin-Elmer instrument. Heated water, flowing through the block can bring the cell temperature to as high as 80°C. Figure 12 shows the phenanthrene excitation and emission spectra taken in air, at 80°C. The concentration

was about 53 ppm, and the spectra are consistent with other measurements (References 2, 7, 8 and 11). The excitation intensity, below approximately 237 nm, should be regarded with suspicion. The xenon lamp intensity falls off dramatically in this wavelength region and there may well be other instrumental artifacts. The feature near 480 nm in the emission spectrum is the second order scatter from the excitation wavelength used (237 nm). Again, it would appear that a 10 ppm level would be the approximate detection limit.

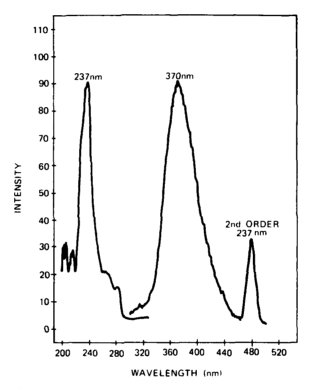


FIGURE 12. Fluorescence Excitation and Emission Spectra of Phenanthrene, Taken in Air, at 80°C.

The fluorescence spectra of pyrene in air, at 80°C, are given in Figure 13. The vapor pressure of pyrene is 2.42 x 10⁻³ at 80°C (Reference 48), which gives a concentration of 3.5 ppm. The greater sensitivity for this PAH follows from the fact that the quantum yield of pyrene is larger than that of both naphthalene and phenanthrene (by a factor of 1.4 and 2.5, respectively, in solution) (Reference 49). The increase in the emission intensity beyond 460 nm is second order scattering from the excitation used (239 nm). By extrapolation, subppm detection levels would be possible with this material.

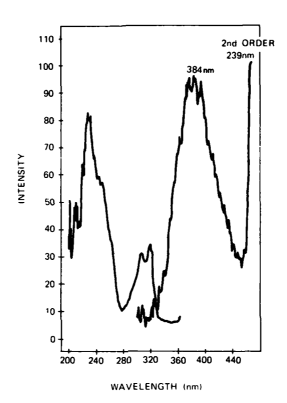


FIGURE 13. Fluorescence Excitation and Emission Spectra of Pyrene, Taken in Air, at 80°C.

The Perkin-Elmer instrument, on which the preceding work was done, is equipped with a 150 W xenon lamp, and its detection electronics work in either a direct current (DC) or an alternating currect (AC) mode. Its response was such that it would not quite meet the 1 ppm PAH detection requirements of the program, and it was felt that further searching was necessary for an appropriate PAH monitor instrument. Photon counting is known to be a more sensitive method of detection than DC or AC amplification. An instrument with photon counting electronics and a greater lamp flux was next considered.

Spex F112 Fluorimeter

After we contacted Spex Industries about their FLUOROLOG 2 series of instruments, they agreed to perform some static cell, gas phase

fluorescence experiments, similar to the ones described in the previous section. The F112 version of their fluorimeter used for these tests had photon counting detection electronics, a 250 nm blazed excitation monochromator grating, and two 500 nm blazed emission monochromator gratings. The correspondence regarding the instrument, tests, and the actual PAH fluorescence spectra are given in Appendix E (Figures A through N).

The gas phase PAH spectra were all measured in air. A cuvette, containing a crystal of the PAH material, was mounted in a water heated brass block to maintain the elevated temperatures. The phenanthrene vapor pressures (Reference 48) at 60, 50, and 30°C are 6.46×10^{-3} , 2.49×10^{-3} , and 3.08×10^{-4} torr, respectively. At sea level (P = 760 torr) this corresponds to 8.50, 3.28, and 0.405 ppm. For pyrene, the corresponding pressures (Reference 48) are 3.64×10^{-4} , 1.29×10^{-4} , and 1.33×10^{-5} torr, which translates into 0.479 and 0.170 ppm, and 17.5 parts per billion (ppb).

The 60°C fluorescence excitation spectrum of pyrene is given in Figure A (Appendix E). The features at 308 and 321 nm compare favorably with Figure 13, but there are some additional features near 265 and 284 nm. The region below 230 nm should again be regarded with suspicion, since instrumental artifacts may be present here, as with the Perkin-Elmer instrument.

The corresponding fluorescence emission spectrum of pyrene is given in Figure B. The sharp feature near 384 nm is due to pyrene (refer to Figure 13), but the broad emission, present from about 410 nm and extending beyond 480 nm, must be an impurity. Nonetheless, the instrument has the capability to detect pyrene at a concentration of 0.479 ppm.

Figures C and D show the fluorescence excitation and emission spectra of phenanthrene at 60°C. They compare quite favorably with Figure 12. Note that phenanthrene shows no absorption (excitation) feature at 321 nm, whereas both phenanthrene and pyrene absorb in the 241 nm region (with phenanthrene dominating). This fact will form the basis for the selective excitation study of a mixture of these two materials (although at least three will be present, given the observation of the impurity in the pyrene sample).

Figures E and F present the fluorescence emission spectra of a pyrene/phenanthrene mixture excited at two different wavelengths. With 241 nm excitation, both molecules absorb, and the phenanthrene emission predominates. With 321 nm excitation, the pyrene (and impurity) emission is selected. These spectra nicely show the advantages of selective excitation for the analysis of multicomponent mixtures.

Figures G and H are a repeat of the spectra of Figures E and F, but the spectra were collected at 50°C . It is important to note that one can still discern the pyrene feature near 384 nm. This is at a concentration of 0.170 ppm. Figures I and J depict the same kind of experiment but at 30°C . The phenanthrene emission can still be seen in Figure I at 0.405 ppm, but only the impurity emission can be seen in Figure J. This is not surprising since the pyrene concentration would be 17 ppb at this temperature.

After discussing these spectra with Spex Industries personnel, particularly concerning the pyrene impurity, they agreed to repeat the pyrene experiments with a fresh sample. The results are given in Figures K through N. Of particular note is the lack of the strong 470 nm impurity emission in Figure K. Both Figures K and N agree reasonably well with Figure 13, ignoring the bluest region of Figure N (especially since this is an uncorrected excitation spectrum).

Comparing Figures F and H, it can be seen that the 384 nm pyrene emission intensity scales well with the concentration change calculated from the vapor pressures. The agreement is not as good in the case of phenanthrene (refer to Figures E, G, and I). Since we have confidence in the vapor pressure values (they were extrapolated from experimental measurements (Reference 48) over the temperature range of 36.70 to 49.65°C), it may have been that the sample was not allowed to reach equilibrium temperature before the fluorescence measurement was Since the pyrene measurements were made following the phenanthrene ones, the sample would have had more time to reach equilibrium. Hence, the better agreement for pyrene. Despite this discrepancy, the Spex F112 fluorimeter is capable of detecting sub-ppm levels of gas phase pyrene and phenanthrene in air. With regard to other PAHs, it should be noted that a number of them have quantum yields greater than these two molecules. For example, benzo(a)pyrene (BaP) has a quantum yield that is about two times greater than pyrene (References 5 and 49). Therefore, one might expect a detection limit for BaP that is about two times lower than for pyrene.

The decision was made to purchase a Spex F112 fluorimeter as the heart of the PAH monitor system, and make the minor flow cell modifications described in the Experimental section.

LANL INCINERATION TESTS

Since we had no previous "hands-on" experience with the Spex fluorimeter prior to our arrival at the LANL, it was decided that it would be best to proceed in stages to get the PAH monitor into operation. Static cell, gas phase experiments were performed first, followed by the flow and calibration modifications to the system. This preliminary work eventually led to the real-time, on-line stack gas sampling

analysis during the Navy colored smoke incineration tests. Because of the time frames involved (the basic instrument was being installed during the incinerator warm-up), some of the minor problems encountered were not dealt with in the most elegant manner, but solved "on the fly" so as to get the instrument on-line during the test schedule. Further system improvements are certainly possible, and a number of suggestions are listed in the Conclusions and Recommendations section.

All the fluorescence spectra were taken with an approximate 5 nm resolution for both the excitation and emission. The wavelengths were scanned in a burst mode at 0.5 nm/step with an integration time of 1 second. A 100 nm spectrum could be obtained in 3.3 minutes at this scan rate. The intensity versus time plots, at fixed excitation and emission wavelengths, were taken with an integration time of 1 second.

In the case of the static cell and calibration flow system measurements, the concentration calculations (equation 3) were done with a value of 585 torr for the atmospheric pressure. The pressure inside the CAI is 492 torr, and this value should be used for all the on-line monitoring situations.

Preliminary Observations

PAH Analysis in a Static Cell. Following the initial installation of the basic Spex F112 fluorimeter, the first experiments were performed with naphthalene in a static cell. The capped sample cuvette (1.0 cm pathlength) was mounted in the Spex model 1931A heater/cooler The block had been modified for temperature control, as described in the Experimental section. Figures 14 and 15 show the naphthalene fluorescence emission and excitation spectra, taken in a front face (FF) emission collection geometry. These spectra agree well with Figure 11, but there was concern about the large background signal level. Unwanted scattering is usually more pronounced in a FF type experiment. To reduce this contribution, and to ensure that only gas phase emission was contributing to the spectra, the emission collection geometry was switched to right angle (RA). The resulting emission spectrum is presented in Figure 16. Note the dramatically reduced baseline, as compared to that in the previous spectra. Based on these results, it was decided to maintain a RA emission collection geometry for all successive spectra.

Next, the solid naphthalene was dumped from the sample cell, and the cell was washed, dried, and replaced in the instrument after a crystal of anthracene was added to it. Upon reexamination, it was found that there was still a small amount of naphthalene in the cell, despite our cleaning efforts. Figure 17 shows the naphthalene emission in the naphthalene/anthracene mixture at 30 and 50°C. The

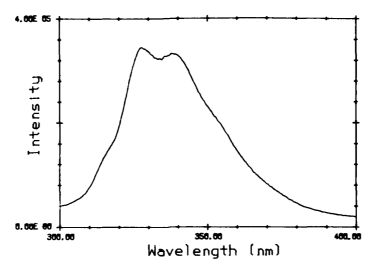


FIGURE 14. Fluorescence Emission Spectrum of Naphthalene in a Static Cell. The temperature was 30°C and the concentration was 239 ppm. A front face emission collection geometry was used. $\lambda_{\rm ex}$ = 268 nm.

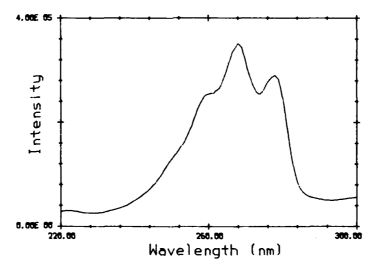


FIGURE 15. Fluorescence Emission Spectrum of Naphthalene in a Static Cell. The temperature was 30°C and the concentration was 239 ppm. A front face emission collection geometry was used. $\lambda_{\rm ex}$ = 328.5 nm.

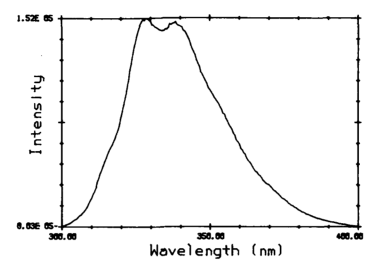


FIGURE 16. Fluorescence Emission Spectrum of Naphthalene in a Static Cell. The temperature was 30°C and the concentration was 239 ppm. A front face emission collection geometry was used. $\lambda_{\rm ex}$ = 268 nm.

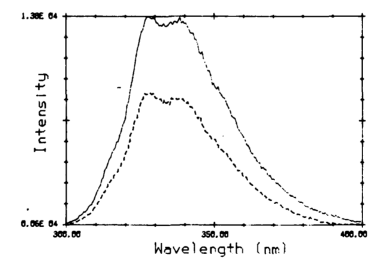


FIGURE 17. Fluorescence Emission Spectrum of a Naphthalene-Anthracene Mixture in a Static Cell. $\lambda_{\rm ex}$ = 268 nm. Comparison at 30°C (---) and 50°C (---).

naphthalene vapor pressure changes by approximately a factor of 8 over this temperature range, but the fluorescence intensity only changed by 1.6. This is because there was not enough naphthalene in the cell to maintain the vapor pressure. A rough calculation, using the ideal gas law, shows that 20 μg would be needed to attain the required 0.809 torr in the 4 ml cell. Although this is enough material to see with the naked eye, we did not see any during the visual inspection of the cell.

The fluorescence excitation and emission spectra of the anthracene in the mixture are presented in Figures 18 and 19. At 50° C, the vapor pressure is 2.13 x 10^{-3} , which corresponds to 3.6 ppm. Figure 19 compares quite well with the results in Appendix A, and other work (References 2, 7, 8 and 11). Figures 17 through 19 clearly show the advantage of selective excitation in a multicomponent mixture analysis. The spectra can be completely separated for the two materials!

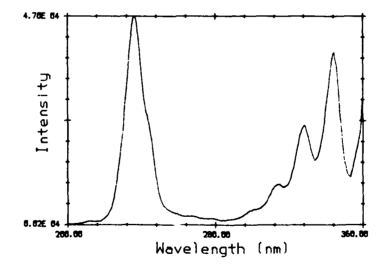


FIGURE 18. Fluorescence Excitation Spectrum of a Naphthalene-Anthracene Mixture in a Static Cell. $\lambda_{\rm em} = 384.5$ nm. The temperature was 50°C and the concentration of anthracene was 3.6 ppm.

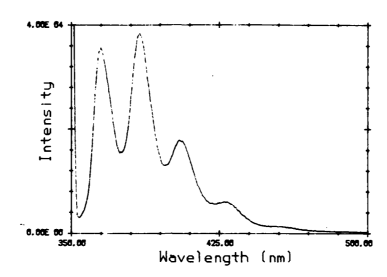


FIGURE 19. Fluorescence Emission Spectrum of a Naphthalene-Anthracene Mixture in a Static Cell. $\lambda_{\rm ex} = 344$ nm. The temperature was 50°C and the concentration of anthracene was 3.6 ppm.

To prove that the fluorescence emission was not coming from solid anthracene that might have been plated-out on the cell windows, the emission spectrum of anthracene crystals is given in Figure 20. It was obtained under 365 nm mercury lamp excitation, which is a wavelength where the gas phase anthracene happens to emit. Comparing Figures 19 and 20, note that there is a completely different structure and wavelength dependence for the emission of the gas and solid phase material. Similar arguments should also hold for the other PAH compounds.

Figures 21 and 22 show the effects of selective excitation on the naphthalene/anthracene mixture for the excitation region near 250 nm. In this case, relatively small excitation wavelength changes can produce either an anthracene, or naphthalene, dominated emission spectrum. The vibronic structure in the anthracene emission (compare Figures 19 and 21) appears washed out when deep UV excitation is used. We are not sure of the reason for this, but an energy transfer process involving the naphthalene may be occurring (References 2, 34 and 25).

To carry the mixture analysis one step further, phenanthrene was added to the cell containing the naphthalene/anthracene. The fluorescence excitation spectrum of this mixture is given in Figure 23. The concentrations for anthracene and phenanthrene were 3.6 and 4.3 ppm. The concentration of naphthalene was probably <30 ppm. This spectrum

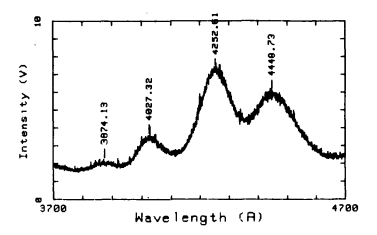


FIGURE 20. Fluorescence Emission Spectrum of Anthracene Crystals. $\lambda_{\rm ex}$ = 365 nm.

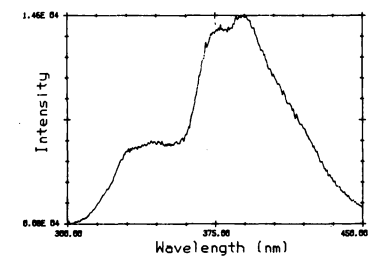


FIGURE 21. Fluorescence Emission Spectrum of a Naphthalene-Anthracene Mixture in a Static Cell. $\lambda_{\rm ex}$ = 247 nm. The temperature was 50°C.

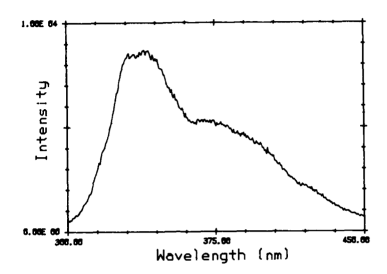


FIGURE 22. Fluorescence Emission Spectrum of a Naphthalene-Anthracene Mixture in a Static Cell. $\lambda_{\rm ex}$ = 251 nm. The temperature was 50°C.

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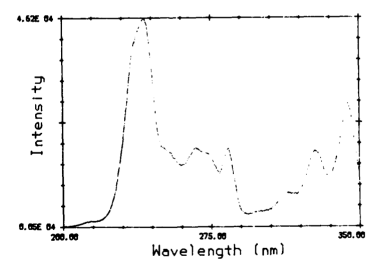


FIGURE 23. Fluorescence Excitation Spectrum of a Naphthalene-Anthracene-Phenanthrene Mixture in a Static Cell. $\lambda_{\rm em}=367.5$ nm. The temperature was 50°C and the concentrations of anthracene and phenanthrene were 3.6 and 4.3 ppm, respectively.

should be compared to the excitation spectra for anthracene (Figure 18) and phenanthrene (Figure D in Appendix E). In fact, it is approximately the addition of the two separate spectra, plus a small contribution from the naphthalene, near 268 nm. When the fluorescence emission is monitored at 367.5 nm, anthracene and phenanthrene contribute strongly (refer to Figure 19 and Figure C of Appendix E). naphthalene emission contribution at this wavelength is somewhat weaker (see Figure 16). The emission spectrum of this mixture, taken with 250 nm excitation, is presented in Figure 24. This excitation wavelength discriminates against anthracene. The strong 370 nm emission is characteristic of phenanthrene (refer to Figure C of Appendix E), and the naphthalene contribution can be seen as a shoulder near 330 nm. There is also an additional feature at 415 nm which must be from an impurity in the phenanthrene.* This impurity contribution to the total (now at least 4-component) emission can be selected with 290 nm excitation, as depicted in Figure 25. This excitation wavelength discriminates against the phenanthrene, anthracene, and naphthalene in the sample. Figures 26 and 27 show the emission spectra of the mixture, irradiated with 284 and 344 nm excitation. The latter is primarily an anthracene emission spectrum.

Using a program provided with the instrument (Spex Tech. Note #62), the computer controlled acquisition of an excitation-emission matrix was performed on the naphthalene/anthracene/phenanthrene/impurity mixture. The result is plotted in Figure 28. It is a series of emission spectra, taken at a series of excitation wavelengths. The vertical axis is intensity. The largest peak corresponds to phenanthrene and anthracene, and one can just make out the naphthalene contribution in the emission scans which were taken with the excitation wavelength at 255 and 260 nm. It took approximately an hour to generate the plot, but it should be empasized that routine work would not need to be done with such a fine grid of points, and could be accomplished much faster. The graph is presented here, mainly to show the type of result that can be obtained quite easily with the Spex instrument.

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In a final prelude to the conversion of the system to flow cell operation, the 33.3°C fluorescence excitation and emission spectra of benzene (another environmentally hazardous material, in its own right) was measured using another applications program (Spex Tech. Note #60). The results are presented in Figure 29. As can be seen from the figure, this program automatically plots the excitation and emission

From the data in Table 3, the measured diffusion rate is approximately 10% higher than the calculated value. The impurity in the phenanthrene could well be a contributing factor to this discrepancy.

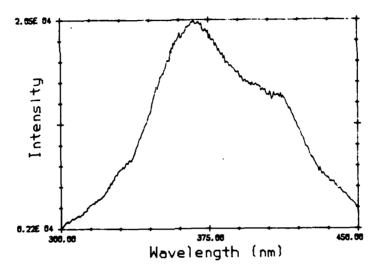


FIGURE 24. Fluorescence Emission Spectrum of a Naphthalene-Anthracene-Phenanthrene Mixture in a Static Cell. $\lambda_{\rm ex}$ = 250 nm. The temperature was 50°C.

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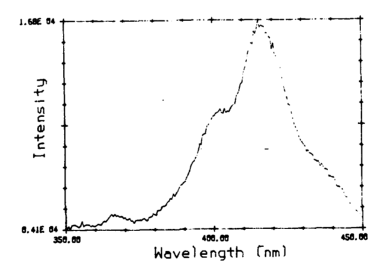


FIGURE 25. Fluorescence Emission Spectrum of a Naphthalene-Anthracene-Phenanthrene Mixture in a Static Cell. Impurity emission selected with $\lambda_{\rm ex}$ = 290 nm. The temperature was 50°C.

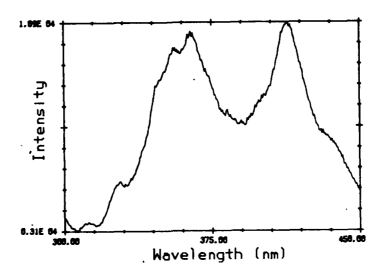


FIGURE 26. Fluorescence Emission Spectrum of a Naphthalene-Anthracene-Phenanthrene-Impurity Mixture in a Static Cell. $\lambda_{\rm ex}$ = 284 nm. The temperature was 50°C.

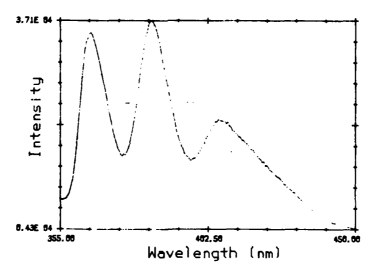


FIGURE 27. Fluorescence Emission Spectrum of a Naphthalene-Anthracene-Phenanthrene-Impurity Mixture in a Static Cell. Selectively excited for anthracene with $\lambda_{\rm ex}=344$ nm. The temperature was 50°C.

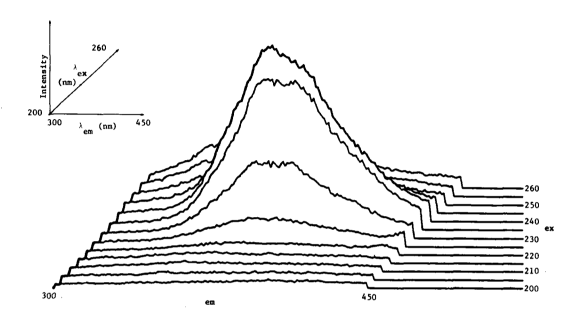


FIGURE 28. Excitation-Emission Matrix Generated from the Naphthalene-Anthracene-Phenanthrene-Impurity Mixture in a Static Cell. The temperature was 50° C. The λ_{ex} values were incremented in 5 nm steps from 200-260 nm. The emission was scanned from 300-450 nm in 1 nm steps. The program was from Spex Technical Note #62.

spectra on the same graph. The gas phase concentration of benzene in the static cell was very high for our purposes (about 200K ppm), but the relevant wavelengths will be useful for the next stage of the work.

PAH Analysis in the Flow System. The instrument was next configured to the PAH monitor flow system of Figures 8 and 9 (described in the Experimental section). Photographs of the actual instrument, at the site, are given in Figures 30 through 32. The diffusion cell, inside the furnace, is shown in Figure 33. The initial flow system experiments were done with benzene. Because of its high vapor pressure (>100 torr at 33.3°C), the operation of the flow system could be checked out with benzene before being heated. Figure 34 shows the instrumental background, taken with the excitation source at the wavelength maximum for benzene (also blue enough to excite other unwanted materials that might be present). At this point, laboratory air from the general instrument location was flowing through the cell. Benzene was next added to the diffusion cell and the fluorescence excitation

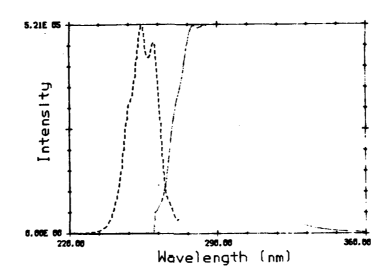


FIGURE 29. Excitation (---) and Emission (--) Spectra of Benzene Vapor in a Static Cell. The temperature was 33.3°C and the concentration was > 200K ppm. The program used to generate the plot was from Spex Technical Note #60.



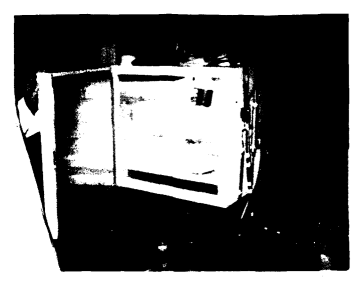
FIGURE 30. PAH Monitor and Flow System Showing Sampling Line to Incinerator.



FIGURE 31. PAH Monitor and Flow System Showing DATAMATE Computer and Operator.



FIGURE 32. PAH Monitor and Flow System.



 $\begin{array}{lll} \textbf{FIGURE 33.} & \textbf{Calibration Cell and Furnace} \\ \textbf{Assembly.} \end{array}$

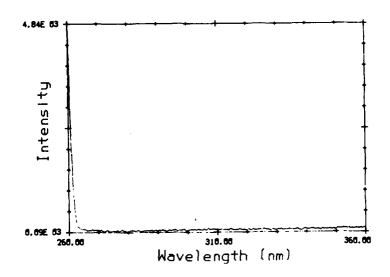


FIGURE 34. Flow Cell Background Spectrum. Excitation and emission parameters selected to detect benzene.

and emission spectra were obtained at several flow rates. Figures 35 through 38 show the excellent agreement between the flow rates (concentrations)* and the observed intensities. The benzene concentrations were high (a few hundred ppm, even at the highest flow rate), and this lack of sensitivity for benzene can be quite easily explained. First, the extinction coefficient for benzene (Reference 49) is 200 L/(mole-cm) near 254 nm, compared to naphthalene's 6,000 L/(mode-cm) at 268 nm. The quantum yield for benzene is also 3.3 times smaller than the yield of naphthalene (Reference 49). This accounts for a two order of magnitude difference in emission intensity between these two materials, not even taking into consideration the fact that the benzene fluorescence may be more highly quenched by air.

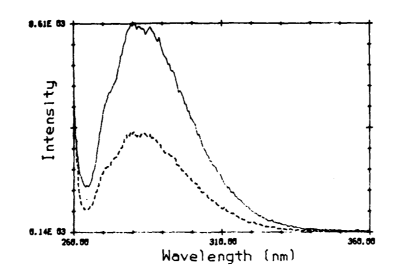
The next day, the cell background was remeasured to be sure that all of the benzene was out of the system. It is shown in Figure 39. Naphthalene was then added to the diffusion cell, and the system was heated to 65.5°C. The transfer lines were normally kept 5 to 10°C higher than the diffusion cell furnace and fluorescence flow cell in order to keep material from condensing in the lines. The fluorescence emission and excitation spectra of naphthalene in the flow system are given in Figures 40 through 43. The DIFF RATE calculations for the 3.9 and 2.3 ppm concentration values are given in Appendix C. It should be noted that the observed intensities again scale quite well with the flow rates (concentrations). The 2.3 ppm plots have been mathematically smoothed. This is part of the standard Spex DATAMATE processing (see Appendix B).

The above experiments indicated that the PAH monitor flow system was in working order, and that the monitor was responding to PAH concentrations on the order of 1 ppm. At this point, the final stage of the work, on-line testing, was begun.

Real-Time Analysis of Navy Colored Smoke

The stack gas temperature at the PAH monitor sampling location (refer to Figures 5 and 6) was 65.5°C during all the incineration tests. The fluorescence cell was also maintained at that set point. The sampling lines were kept 5 to 10°C higher than the sampling location temperature in order to prevent the possibility of condensation in the lines. Because of the low sampling temperature, the PAH monitoring wavelengths were chosen to detect the smaller, higher vapor pressure materials. The larger materials, if present, would most likely be

Although the flowmeter curves were calibrated for air at 760 torr and 70°F, the calculated correction factors for the LANL altitude and temperature were found to be generally less than 10%. In what follows, no correction was used.



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FIGURE 35. Fluorescence Emission Spectrum for Benzene Vapor in the Flow System. $\lambda_{\rm ex}=254$ nm. The temperature was 33.3°C and the flow rates were 22 mL/min (---) and 44 mL/min (---).

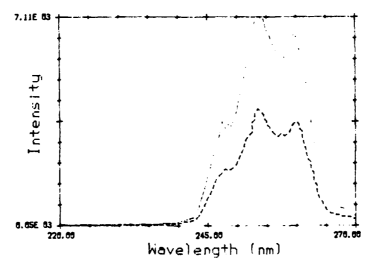


FIGURE 36. Fluorescence Excitation Spectrum for Benzene Vapor in the Flow System. λ = 285 nm. The temperature was 33.3°C and the flow rates were 22 mL/min (---) and 44 mL/min (---).

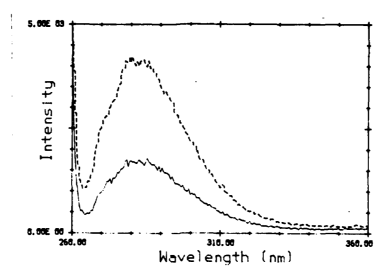


FIGURE 37. Fluorescence Emission Spectrum for Benzene Vapor in the Flow System. $\lambda_{\rm ex} = 254$ nm. The temperature was 33.3°C and the flow Fates were 110 mL/min (---) and 44 mL/min (---).

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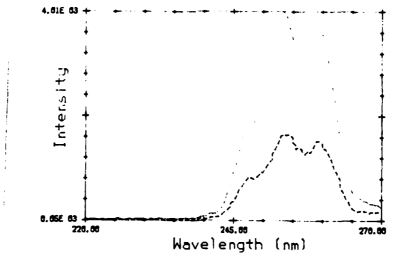


FIGURE 38. Fluorescence Excitation Spectrum for Benzene Vapor in the Flow System. $\lambda_{\rm em} = 285$ nm. The temperature was 33.3°C and the flow rates were 44 mL/min (---) and 110 mL/min (---).

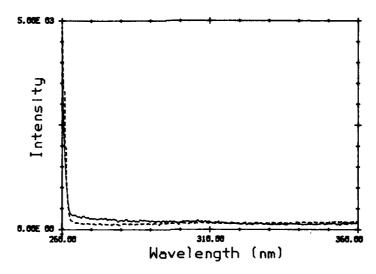


FIGURE 39. Comparison of the Flow Cell Background Spectrum for Two Separate Days. Figure 34 is the second spectrum (---). Excitation and emission parameters were selected to detect benzene.

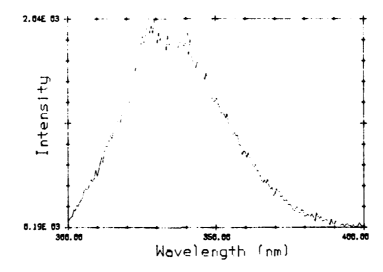


FIGURE 40. Fluorescence Emission Spectrum for Naphthalene in the Flow System. $\lambda_{\rm ex}=269$ nm. The temperature was 65.5°C and the flow rate was 160 mL/min. The concentration of naphthalene was 3.9 ppm.

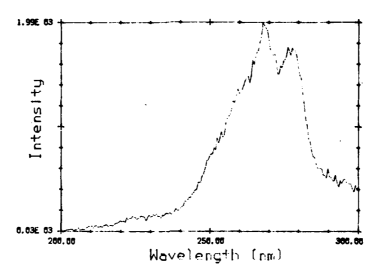


FIGURE 41. Fluorescence Excitation Spectrum for Naphthalene in the Flow System. $\lambda = 328$ nm. The temperature was 65.5°C and the flow rate was 160 mL/min. The concentration of naphthalene was 3.9 ppm.

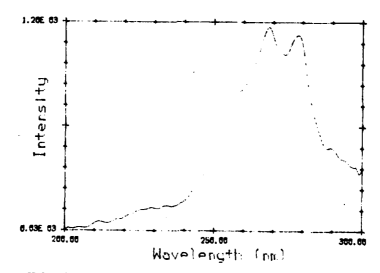


FIGURE 42. Fluorescence Excitation Spectrum for Naphthalene in the Flow System. λ = 328 nm. The temperature was 65.5°C and the Tlow rate was >270 mL/min. The concentration of naphthalene was 2.3 ppm.

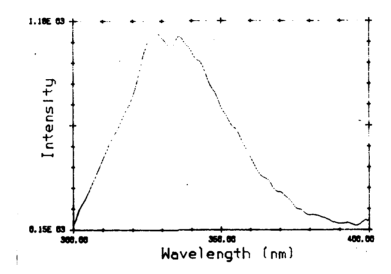


FIGURE 43. Fluorescence Emission Spectrum for Naphthalene in the Flow System. $\lambda_{\rm ex} = 269$ nm. The temperature was 65.5°C and the flow rate was \geq 270 mL/min. The concentration of naphthalene was 2.3 ppm.

condensed on particulates. For example, BaP has a vapor pressure of 5.49 x 10⁻⁹ torr at 25°C (Reference 2). Even allowing for a factor of 20 increase at 65.5°C, the ambient concentration would be approximately 0.2 ppb. This is well below the detection capabilities of the monitor as configured and operated here. If, for some reason larger PAHs were present, they would be excited by the UV wavelengths used for the smaller compounds, and would contribute to the emission at redder wavelengths. Therefore, their possible presence was not entirely ignored in the work. The actual chemical composition of the Navy colored smokes have been defined previously (Reference 1), and the CAI feed schedules and operating conditions are given in the LANL experimental test plan (Reference 41).

The results from the first attempt to put the PAH monitor system on-line are given in Figure 44. The Mk 13 colored smoke composition was being incinerated at the time. Unfortunately, the incineration test was aborted a few minutes after the monitoring began, because the incinerator was running at the wrong temperature for the Mk 13, phase 4, period 1 test. The gas sampling was initiated at t = 116 seconds on the plot, and the excitation and emission wavelengths of the instrument were set for the naphthalene maxima. The important thing to

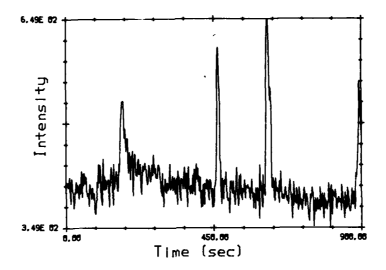


FIGURE 44. On-Line Temporal Scan Started at 1204 Hours on 11 September 1983. Mk 13 smoke composition. λ = 269 nm, λ = 328 nm. Wavelength parameters for naphthalene detection.

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note is that after a momentary response at the switching point no significant change occurred in the background level. There also were a few additional intensity spikes in the 450 to 900 second region. At this time, their origin is uncertain. It is tempting to speculate that these spikes may have been caused by the instrument response to PAHs deposited on particulate matter, but given that the sampling location was situated between two HEPA filters, this is unlikely.

While the CAI conditions were being reset for the next incineration test, the PAH monitor was switched back to the calibration cell (containing naphthalene). The time response of the system was next examined by monitoring the fluorescence intensity as the diffusion cell sample tube was replaced with a clean, empty plug. Figure 45 shows the decay of the naphthalene signal after the insertion of the empty plug (at t = 80 seconds). A steady state is reached in a few minutes.

The monitor system was next switched back on-line for the Mk 13, phase 4, period 1 test as shown in Figure 46. Again, the naphthalene excitation and emission wavelengths were used for detection. Of particular significance is the fact that the signal intensity actually

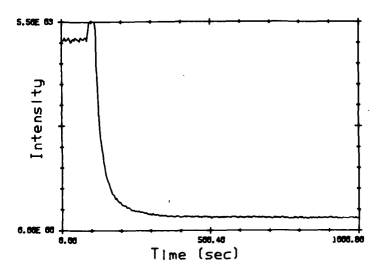


FIGURE 45. Temporal Scan During the Removal of a Naphthalene Calibration Sample and the Insertion of an Empty Plug. $\lambda_{ex} = 269 \text{ nm}$, $\lambda_{em} = 328 \text{ nm}$. Wavelength parameters for naphthalene detection.

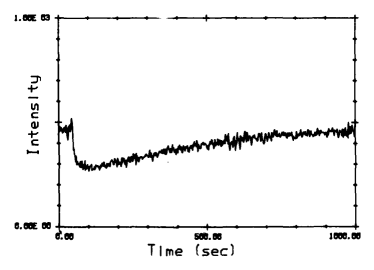


FIGURE 46. On-Line Temporal Scan Started at 1445 Hours on 11 September 1983. Mk 13, phase 4, period 1. λ_{ex} = 269 nm, λ_{em} = 328 nm. Wavelength parameters for naphthalene detection.

decreased when the instrument was put on-line (it was switched over from the empty diffusion cell configuration). This may mean that the incinerator gases at the sampling point were actually cleaner than the laboratory air flowing through the empty diffusion cell. There was, in fact, a distinct fuel-oil odor in the room at this time. The lower pressure (492 torr) inside the incinerator system, relative to the ambient atmospheric pressure (585 torr) may be a contributor to the signal decrease. Figures 47 through 52 are a series of spectra taken during the Mk 13, phase 4, period 1 incineration test. The acquisition times are given on the graph labels in 24-hour clock notation. In all cases, the signals were not caused by the presence of PAHs on a ppm level, but caused by either a background from the flow system, or scattered light contributions.* Figure 53 was generated during the switch from off-line, to the empty diffusion cell configuration (at t = 320 seconds).

Figures 54 through 60 were all taken during the Mk 13, phase 4, period 2 test. The CAI burner went out during the scan of Figure 56 (at t = 1925 hours), and Figure 57 was acquired, on-line, but with the burner down. The system was switched to the empty calibration cell for Figure 58, which shows the system background, and Figures 59 and 60 were generated with the burner functioning again. No PAHs were detected on a ppm level for this entire test.

Similar negative results (no PAHs) are shown in Figures 61 through 64, measured during the Mk 21, phase 6, period 1 incineration test. Again, the signal level decreased when the system was put on line.

Figure 65 shows the negative results for the Mk 89, phase 7, period 1 test, and Figure 66 is an excitation-emission matrix, generated while on-line. It took approximately 25 minutes to generate. Again, routine analysis would not need to be done with such a fine grid of data points, and the task in survey fashion could easily be accomplished more rapidly.

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A final negative test result is shown in Figure 67. After this run, the system was switched back to the empty diffusion cell sampling configuration. Naphthalene was then put into the diffusion cell sample tube, and the time response of the system was measured. As can be seen from Figures 68 and 69, it took about an hour to reach the steady

[&]quot;It was noticed that there was scattered light coming from the insulating tape used on the inlet flow line to the fluorescence cell. Baffling material was inserted into the sample compartment to temporarily improve the situation, but this problem should be more permanently addressed in future work.

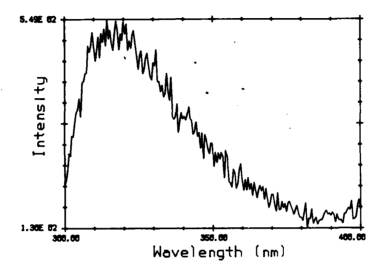


FIGURE 47. On-Line Emission Scan Started at 1500 Hours on 11 September 1983. Mk 13, phase 4, period 1. $\lambda_{\rm ex} = 269$ nm. Wavelength parameters for naphthalene detection.

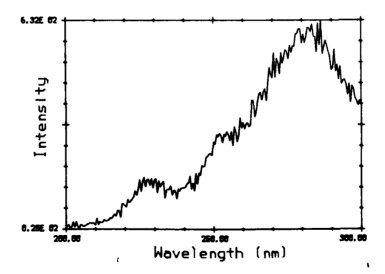


FIGURE 48. On-Line Excitation Scan Started at 1510 Hours on 11 September 1983. Mk 13, phase 4, period 1. $\lambda_{em} = 328$ nm. Wavelength parameters for naphthalene detection.

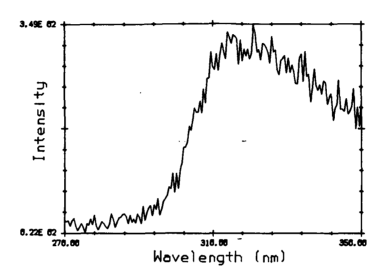


FIGURE 49. On-Line Emission Scan Started at 1517 Hours on 11 September 1983. Mk 13, phase 4, period 1. $\lambda_{\rm ex}$ = 254 nm. Wavelength parameters for benzene detection.

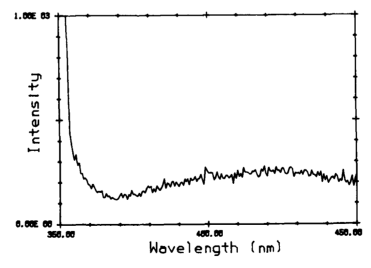


FIGURE 50. On-Line Emission Scan Started at 1530 Hours on 11 September 1983. Mk 13, phase 4, period 1. λ = 344 nm. Wavelength parameters for anthracene detection.

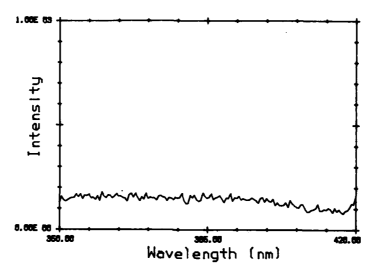


FIGURE 51. On-Line Excitation Scan Started at 1642 Hours on 11 September 1983. Mk 13, phase 4, period 1. $\lambda_{em} = 440 \text{ nm}$.

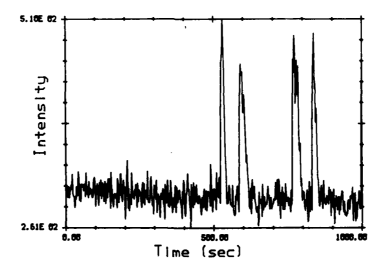


FIGURE 52. Temporal Scan Started at 1652 Hours on 11 September 1983. Just at the end of the Mk 13, phase 4, period 1 run. λ = 269 nm, λ = 328 nm. Wavelength parameters for naphthalene detection.

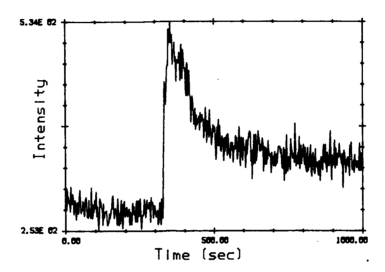


FIGURE 53. Temporal Scan Started at 1715 Hours on 11 September 1983. Switched form on-line to empty diffusion cell (room air) at t = 320 seconds. λ = 269 nm, λ = 328 nm. Wavelength parameters for naphthalene detection.

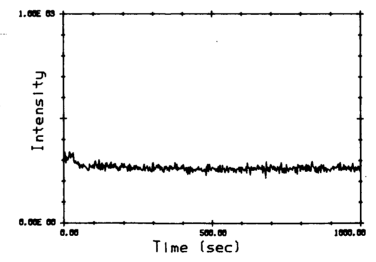


FIGURE 54. Temporal Scan Started at 1843 Hours on 11 September 1983. Switched from empty diffusion cell (room air) to on-line at t = 40 seconds. $\lambda = 269 \text{ nm}, \lambda = 328 \text{ nm}.$ Wavelength parameters for naphthalene detection. Just at start of Mk 13, phase 4, period 2.

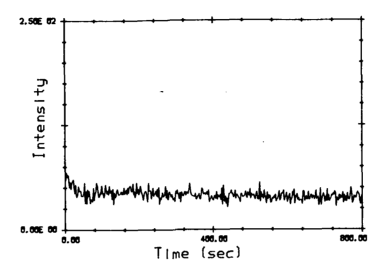


FIGURE 55. On-Line Temporal Scan Started at 1906 Hours on 11 September 1983. Mk 13, phase 4, period 2. λ_{ex} = 254 nm, λ_{em} = 285 nm. Wavelength parameters for benzene detection.

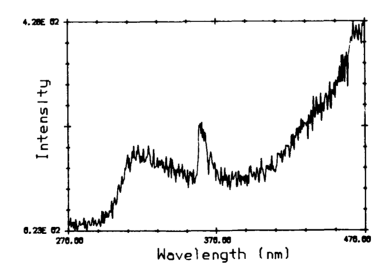


FIGURE 56. On-Line Emission Scan Started at 1924 Hours on 11 September 1983. Mk 13, phase 4, period 2. $\lambda_{\rm ex}$ = 254 nm. Wavelength parameters for benzene detection.

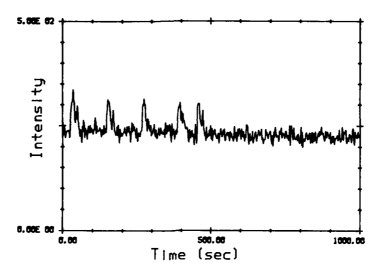


FIGURE 57. On-Line Temporal Scan Started at 1939 Hours on 11 September 1983. Mk 13, phase 4, period 2. $\lambda_{ex}=254$ nm, $\lambda_{em}=328$ nm. The CAI burner went out during this scan.

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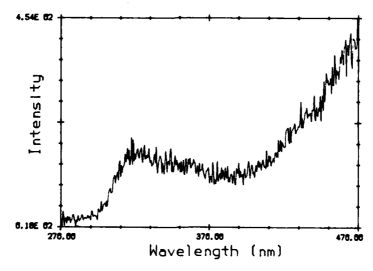


FIGURE 58. Empty Diffusion Cell (Room Air) Background Emission Scan Started at 2015 Hours on 11 September 1983. λ = 254 nm. Wavelength parameters for benzene detection.

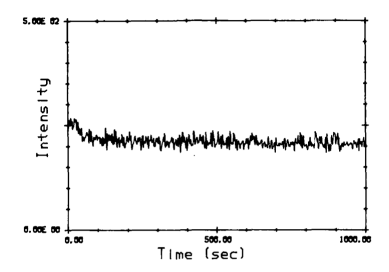


FIGURE 59. On-Line Temporal Scan Started at 2052 Hours on 11 September 1983. Mk 13, phase 4, period 2. $\lambda_{ex} = 269 \text{ nm}$, $\lambda_{ex} = 328 \text{ nm}$. Wavelength parameters for naphthalene detection. The CAI burner was functioning during this scan.

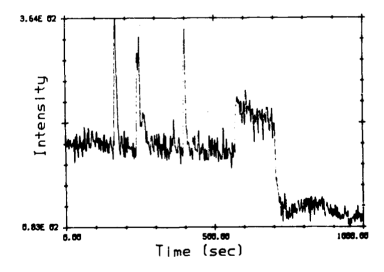


FIGURE 60. Temporal Scan Started at 2110 Hours on 11 September 1983. Just at the end of the Mk 13, phase 4, period 2 run. $\lambda_{\rm ex} = 269$ nm, $\lambda_{\rm ex} = 328$ nm. Wavelength parameters for naphthalene detection.

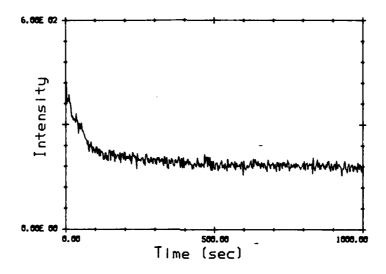


FIGURE 61. On-Line Temporal Scan Started at 1150 Hours on 12 September 1983. Mk 21, phase 6, period 1. $\lambda_{\rm ex} = 269$ nm, $\lambda_{\rm em} = 328$ nm. Wavelength parameters for naphthalene detection.

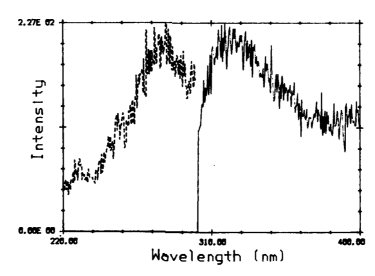


FIGURE 62. On-Line Excitation (---) and Emission Spectra Taken at 1230 Hours on 12 September 1983. Mk 21, phase 6, period 1. The program used to generate the plot was from Spex Technical Note #60.

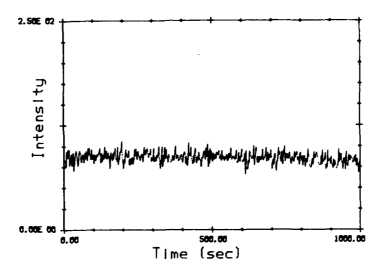


FIGURE 63. On-Line Temporal Scan Started at 1252 Hours on 12 September 1983. Mk 21, phase 6, period 1. $\lambda_{\rm ex}$ = 240 nm, $\lambda_{\rm em}$ = 385 nm.

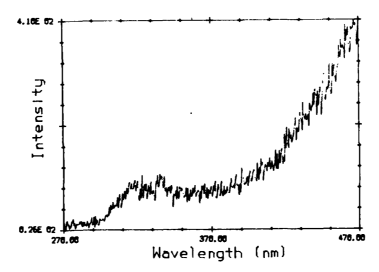


FIGURE 64. Fluorescence Emission Scan Started at 1319 Hours on 12 September 1983. Just at the end of the Mk 21, phase 6, period 1 run. $\lambda_{\rm ex}$ = 254 nm.

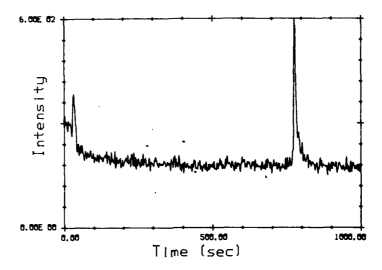
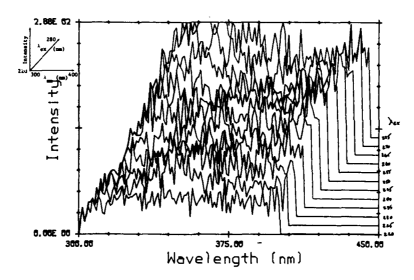


FIGURE 65. On-Line Temporal Scan Started at 0833 Hours on 13 September 1983. Mk 89, phase 7, period 1. $\lambda_{\rm ex} = 269$, $\lambda_{\rm em} = 328$ nm. Wavelength parameters for naphthalene detection.



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FIGURE 66. On-Line Excitation-Emission Matrix Generated at 0900 Hours on 13 September 1983. Mk 89, phase 7, period 1. The λ values were incremented in 5 nm steps from 220-275 nm. **The emission was scanned from 300-400 nm in 1 nm steps. The program was from Spex Technical Note #62.

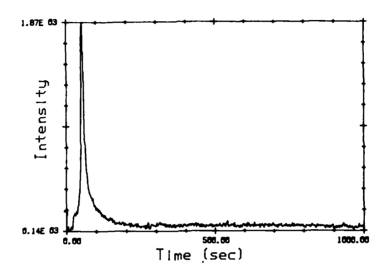


FIGURE 67. Temporal Scan Started at 1814 Hours on 14 September 1983. Taken after the Mk 23, phase 10, period 1 run. λ = 269 nm, λ = 328 nm. Wavelength parameters for naphthalene detection.

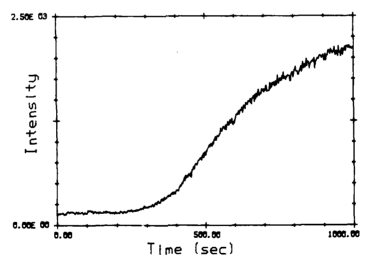


FIGURE 68. Temporal Scan Taken During the Insertion of Naphthalene into the Calibration Cell. The temperature was 65.5° C and the flow rate was 80 mL/min. $\lambda_{ex} = 269$ nm, $\lambda_{ex} = 328$ nm. Wavelength parameters for naphthalene detection. Upon reaching steady state, the naphthalene concentration would be 7.8 ppm.

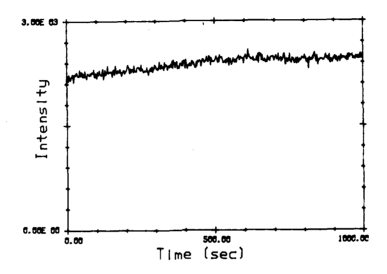


FIGURE 69. Continuation of Temporal Scan of Figure 68.

state. This is quite different from the result given in Figure 45, or the time to steady state (Reference 36) calculations that are given in Appendix C. The causes for this include the fact that the furnace cools down when the door is opened for the insertion of the naphthalene sample and that the room temperature naphthalene is held in an insulating glass sample tube. This time lag is important only when the sample is initially placed in the furnace. The times required for equilibration during flow rate changes occur on a scale more in line with the calculated time to steady state values.

Finally, an excitation-emission matrix was generated for the naphthalene sample. It is given in Figure 70, and should be compared with the on-line test data of Figure 66. The difference between the two situations is quite clear from the figures.

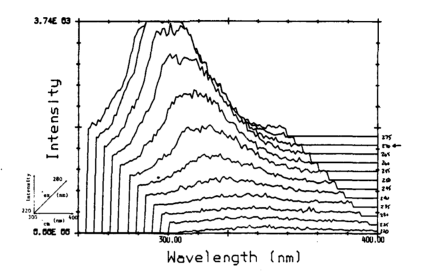


FIGURE 70. Excitation-Emission Matrix Generated from 7.8 ppm Naphthalene. The λ_{ex} values were incremented in 5 nm steps from 220-275 nm. The emission was scanned from 300-400 nm in 1 nm steps. The program was from Spex Technical Note #62.

CONCLUSIONS AND RECOMMENDATIONS

CONCLUSIONS

A number of Navy colored smoke compositions were incineration tested at the LANL CAI facility. A gas monitor analyzed the incinerator effluent for environmentally hazardous PAH materials during the tests. At the sampling location, no PAHs were detected (on a ≥ 1 ppm level) during the incineration of any of the smoke compositions studied. This was true for all the incinerator conditions used over approximately 7 days of testing.

The real-time PAH monitor system, the design, construction, and operation of which is the main focus of this report, is based on the minor modification of a commercially available, computer controlled spectrofluorimeter. The instrument is modular in design, can be upgraded, and has the advantage of being an off-the-shelf item. It performed admirably during the incineration tests, especially when one

considers the rather hostile environment in which it was placed. The monitor operated, despite the noise, vibration, dust, and >98°F temperatures at its location in the CAI facility.* It also survived three major power failures during the tests, and while being installed, had some red dye (probably from Mk 13) settle-out on it.

The computer system for spectrometer control, data manipulation, and spectral storage is user-friendly, and powerful data acquisition programs can be easily written. A number of these are supplied with the basic instrument.

The diffusion-based calibration cell was constructed and installed in the flow system to the PAH monitor. Laboratory testing demonstrated that this device was capable of producing defined concentrations of PAH compounds. This provided a method to check the monitor response to the PAHs. The known concentrations that were generated with the diffusion cell can also be calculated to within 10% accuracy, given the diffusion cell geometry, the vapor pressures of the PAHs used, and the gas flow rates as parameters.

A standard operating procedure (SOP) for the maintenance and automatic operation of the PAH monitor system will ultimately be prepared. The SOP will most likely be based on the excitation-emission matrix idea (References 30 through 33) but with a lower density of points than that obtained with video fluorimetry (References 30 and 31). Finally, the real-time results, presented here, will be compared with the analytical results from the collected samples when those data become available.

RECOMMENDATIONS

Listed below are a number of recommendations meant to improve the PAH monitor system, or its operation, for future work:

- 1. The temperature controller for the cartridge heaters in the fluorescence flow cell block is of the on/off type. This leads to some degree of overshoot in temperature when the controller comes on. A proportional heater/controller would give a more constant temperature set point.
- 2. Spex Industries sells a set of four-fused silica windows that would thermally isolate the sample compartment from the rest of the

Spex Industries claims that the disk drives on the Datamate are prone to read/write errors at >125°F, and that the keyboard keys soften at >160°F.

instrument (about \$160.00 for the set of 4). These would keep the spectrometers in better calibration, especially if higher operating temperatures were used.

- 3. A line needs to be run from the diffusion cell inlet flowmeter to the outside of the building so that "clean" air can be used for calibration purposes. Laboratory air from near the furnace was being used during these tests.
- 4. The spectral data should have been gathered in a corrected, ratioed mode. This is especially true for the excitation spectra, since the lamp intensity is a strong function of wavelength in the deep UV. The spectra reported in this document were not corrected, or ratioed. Therefore, some error was possible due to long term lamp intensity drift. The excitation spectra are also distorted below 240 nm, but since they were all run consistently (i.e., all uncorrected), the results are still valid.
- 5. Permanent baffling should be installed in the sample chamber of the instrument to reduce the unwanted scattering contributions from the insulating tape on the flow cell inlet line. A darker insulating material should also be used.
- 6. Appropriate bandpass glass filters should be used on the emission (and excitation) beam paths so as to further reduce the level of unwanted scattered light.
- 7. If condensation onto the fluorescence cell windows ever becomes a problem, as part of normal operating procedure, the cell temperature could be periodically raised to bake-out the unwanted condensate. (Note that with the simple design of the flow system connections, the flow cell can also be easily removed for cleaning.)
- 8. All the spectra reported here were obtained at 5 nm resolution. Because of the broadband nature of these gas phase spectra, the bandpass of the spectrometers could be increased to 10 nm. This would produce an added factor of $2(2^2) = 8$, in throughput, reducing the detection limits by this same factor.
- 9. Although the flow system, as used in this work, performed reasonably, it should be noted that the switch-over from the calibration cell to the on-line sampling configuration was always done with the diffusion cell empty. This allowed an on-line measurement to be directly compared to the (presumably) null background. If naphthalene were in the diffusion cell at the time, one would be measuring the decay of the naphthalene signal while on-line. This would interfere with the measurement. An additional air line could be added to the system for null measurements, but this would complicate the flow design and the operation of the monitor. For the LANL tests (65.5°C),

removal and insertion of the diffusion cell sample tube represents only a minor annoyance, but for higher temperature operation, this problem would have to be addressed. The question of how often one needs to check the calibration of the instrument should also be asked. If this needs to be done only as often as does the cleaning of the fluorescence flow cell, a simpler configuration of the monitor system might involve the complete removal of the diffusion cell. In this case, a calibration check of the instrument could be done with a static cell, while the flow cell was removed for cleaning.

A few additional operational scenarios that may prove useful in future work are as follows:

- 1. A triangular, quartz flow cuvette (costing a few hundred dollars) would allow this same instrument to be used as an absorption spectrophotometer, if desired. The triangular cuvette would be inserted into the heated brass block such that the excitation beam enters the cell at normal incidence and passes into the sample. The beam then reflects (4 to 8%) off the hypotenuse of the triangle and passes out of the cell, into the emission monochromator. To generate a spectrum, both spectrometers (excitation and emission) would be scanned synchronously.
- 2. The fluorimeter can also be used for routine analysis of collected samples. One need only let the system come to room temperature and remove the flow cell. For example, it may be that as part of the SOP for the Navy incinerator system, an operator must periodically swab samples from the incinerator lines to check for the buildup of dye, or PAH material. These samples can then be extracted and put in a standard cuvette. A fluorescent analysis could be done with no modification of the instrument. Another important point is that the density of a solution sample is approximately one thousand times greater than that of air. The fluorescence quenching would also be less of a factor in solution. This all means that one could approach the advertised 0.1 parts-per-trillion (ppt) detection limit for these samples. Solid filter materials could also be examined, and the use of sensitized-fluorescence techniques (among others) would be possible (References 34, 35 and 50).
- 3. With some loss in the real-time feature, one can insert a trap in the sampling line and concentrate possible PAHs there before analysis. Presumably, one would warm the trap up (with slower flowrate conditions) and subsequently make fluorescence measurements. Another variant of this idea is to insert a gas chromatographic column into the line to aid in the separation of multicomponent mixtures.

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LIST OF COMPANY NAMES

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Beckman Instruments, Inc. Fullerton, CA 92634

CVI Laser Corp.
Albuquerque, NM 87112

Eastman Chemical Products, Inc. Kingsport, TN 37662

Entropy Environmentalists, Inc. Research Triangle Park, NC

Fisher Scientific Co. Pittsburgh, PA 15219

Gilmont Instruments Great Neck, NY 11021

Gow-Mac Instrument Co. Bound Brook, NJ 08805

Hewlett-Packard Co. Westlake Village, CA 91360

Lumonics Inc. Kanata (Ottawa), Ontario Canada K2K1Y3

Matheson Div. Searle Medical Products USA, Inc. E. Rutherford, NJ 07073

Mettler Instrument Corporation Hightstown, NJ 08520

NSG Presicion Cells, Inc. Hicksville, NY 11802

Omega Engineering, Inc. Stamford, CT 06907

Products for Research, Inc. Danvers, MA 01923

Quanta Ray, Inc. Mountain View, CA 94040

RCA Solid State Division Lancaster, PA 17604

Spex Industries, Inc. Edison, NJ 08817

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Appendix A

PROGRAM AD_RLTIME

```
PROGRAM AD_RLTIME AS OF 7/22/83
20
          PROGRAM "AD_RETIME" TO TAKE A/D READINGS IN REAL-TIME.PLO1 DATA! AND CALCULATE STANDARD DEVIATION. ALSO OPTION TO STORE DATA
30
40
50
             ON DISC
60
            USE WITH "DSC RDTM" PROGRAM
80
          IR. LODA 3/17/83
! CURSER INSERTED, P. PLASSMANN 7/20/83
90
100
110
120
130
          ! USE READ/DATA STATEMENTS FOR SELDOM CHANGING VARIABLES-SEE BELOW OPTION BASE 1 !ARRAYS START AT 1 NOT 0 INDEX DIM Seconds(7200).Rav(7200).Std_dev(7200).Ts(81.R(1000)
140
150 DIM Opr$[18],Date$[8],Comment$[40].Pmt$[10].File_name$[10].Dsc_lbl$[6]
160 REAL T_ov.T_dc.T_fc.Flow_f.Flow_g.Tn_cnst.Lam_ex.Bndwth.Power.Re_rate.Lam_e
m.Band.Slits.Hvs.Hvr.Gate_time.Ap_dly.fcs.Tcr.Sens_s.Sens_r
          INTEGER Run_num.N.Test_length.Tm_test
        GOSUB Clear_screen
DUMP DEVICE IS 9
!OUTPUT 9 USING 2100
IMAGE /.30X."START OF EXPERIMENT"
180
190
200
210
220
          PRINTER IS 1
          PRINT TABXY(5,9)."(EXECUTE) SET TIME ENTIME('hh:nm:ss').THEN CONTINUE" PRINT TABXY(5,10)."IF NECESSARY, OTHERWISE JUST CONTINUE."
230
240
250
          PAUSE
        GOSUB Clear_screen
T$=FNT:mes(TIMEDATE)
!INPUT "ENTER OPERATOR NAME(19CHS).DATE(8CHS)".Opr$.Dates
!INPUT "ENTER SAMPLE NAME(18CHS).RUN NUMBER",Sample$.Run_num
!INPUT "ENTER TEMPERATURES-OVEN.DIFFUSION CELL,FLUORESCENCE CELL".T_ov.T_=
260
270
280
290
300
           !INPUT "ENTER GAS NAMES FOR FLOWMETERS F.G(18CHS)".Cas_fs.Gas_gs
!INPUT "ENTER STARTING FLOWRATES-F.G IN CM".Flow_f,Flow_g
320
330
340
350
          READ Uprs.Dates,Samples,Run_num.T_ov.T_dc.T_fc.Gas_fs.Gas_gs.Flow_f.Flow_s
DATA_LODA/PLASSMANN.7/22/83.PHENANTHRENE_N2.1.118.118.118.NITROGEN.LAB_ATE
360
370
         READ N.Tm_cnst.Test_length.Xticno
        DATA 20.1.100,12
390
400
410
420
430
          !INPUT "ENTER N SAMPLES AVERAGED(1000MAX). TIME CONSTANT OF AVERAGE(2<-N<=
! BASED ON .01 SEC READ-TIME
          IF N<=100+Im_cnst THEN
450
460
          GOTO 510
470
          ELSE
          BEEP
480
          GOTO 430
490
          END IF
!INPUT "ENTER TEST LENGTH-SECONDS(7200 MAX).NUMBER OF X TICS ".Test_length
500
510
 .Xticno
           IF Test_length>7200 THEN 510
           IF (Test_length/Tm_cnst<=7200) THEN
```

```
GOTO 720
540
550
          ELSE
560
          Test_length=7200*Tm_cnst
570
580
590
          DISP "MAX TEST LENGTH IS ":Test_length:" FOR A ":Tm_cnst:" TIME CONSTANT."
600
          WAIT 10
510
          BEER
620 INPUT "ENTER O TO CONTINUE WITH THIS TEST LENGTH.ANY KEY TO REENTER TIME C
ONSTANT".Jnk
630 IF Jnk=0 Then
          GOTO 680
540
650
         ELSE
660
670
          GOTO 430
          END IF
          INPUT "NUMBER OF X TICS".Xticno
680
630
700
710
          READ Comments, Sources, Lam_ex, Bndwth, Dyes, Power, Rep_rate, Monos
DATA 75MM LENS-EM-0160 IRTS-EX, EXCIMER (XeCl), 308, 1, NONE, 1, 4, 24, SPEX_.85M
720
730
          READ Lam_em.Band.Slits.PmtS.Hvs.Hvr.Proc_instS.Gate_time.Ap_dly.Tcs.Tcr.Se
740
        Sens_r
DATA 390.0.1.RCA C31034.-1650.-850.DCA-1..1.0.1..3.30.10
75<del>0</del>
760
770
780
790
300
          ! CALC WAIT TIME FOR READ CYCLE
810
820
          Loop_time=.009!NOTE WALTS ARE ROUNDED TO NEAREST THOUSANDTH.
         Wt_time=(Tm_cnst/N)-Loop_time
!INPUT "ENTER COMMENTS(40CHS)".Comments
!INPUT "ENTER EXCITATION SOURCE NAME(18CHS)".Sources
!INPUT "ENTER EXCITATION WAVELENGTH(NM).BANDWIDTH(NM).DYE(18CHS)".Lam_ex.3
830
340
850
860
ndwth.Dyes
          !INPUT "PONER(WATTS).REP_RATE(HZ)".Power.Rep_rate
870
         P_power=(Power/Rep_rate)*1000 !PULSE POWER MJ/PULSE
!INPUT "ENTER MONOCHROMATOR TYPE(18CHS)", Mono$
!INPUT "ENTER EMISSION WAVELENGTH(NM).BANDWIDTH(NM),SLITS(MM)",Lam_em,Banc
880
390
900
.Slits
910
.SITES
910 !INPUT "ENTER PMT TYPE(10CHS).HIGH VOLTAGE-SAMPLE.REFERENCE",Pmt$.Hvs,Hvr
920 !INPUT "ENTER PROCESSING INSTRUMENT(18CHS)",Proc_insts
930 !INPUT "ENTER GATE TIME(MSEC).APERATURE DELAY(MSEC).TIME CONSTANTS-SAMPLE.
REF(SEC)".Gate_time.Ap_dly.Tcs.Tcr
940 !INPUT "ENTER SENSITIVITY OR GAIN-SAMPLE.REF".Sens_s.Sens_r
         GOSUB Clear_screen
GOSUB Data_print
DISP "PAUSE-PRESS CONTINUE TO GO ON"
950
960
970
980
          PAUSE
         GOSUB Clear_screen
PRINTER IS 9
OUTPUT 1:"THERMAL PRINTER ACTIVE"
990
1000
1010
          GOSUB Plot_axes
Tm_test=INI(Test_length/Tm_cnst)
1020
1030
          DISP "PAUSE-PRESS CONTINUE TO START SCAN"
PAUSE
DISP_"SCANNING"
1040
1050
 1060
          GOSUB Start_timer
 1070
1080
```

```
1090
            ! MAIN LOOP
           ON KEY 2 LABEL "FLOW CHANGE" GOSUB Flow_change FOR J=1 TO Tm_test
1100
1110
1120
            Sum=0.
            Sum_sq=0.
GDSUB Read_ad
GDSUB Plot_inten
1130
1140
1150
1160
              NEXT J
1170
1180
            ! END MAIN LOOP
1190 GOSUB Knob_thing
1200 Plot_axes: DEG
                                 GRAPHICS ON
1210
1220
                                 GINIT
1230 LORG 5
1240 READ Label_x$.Label_y$.Vxmin,Vxmax.Vymin.Vymax.Xmin.Ymin.Ymax,Yticno
1250 Xmax=Test_length
1260 DATA TIME-SEC.INTENSITY.20.110.31,90.0.0.10.10
           WINDOW Xmin.Xmax.Ymin.Ymax
Xtic=(Xmax-Xmin)/Xticno
Ytic=(Ymax-Ymin)/Yticno
AXES Xtic.Ytic.Xmin.Ymin.Xticno.Yticno,5
VIEWPORT 0.131.0.100
1280
1290
1300
 1310
 1320
           UDIR 0

MOVE Xmin+(Xmax-Xmin)/2.Ymin-.10+(Ymax-Ymin)
CSIZE 7+(Vxmax-Vxmin)/131..6

LABEL USING "K":Lanel_x$

MOVE Xmin-.10+(Xmax-Xmin).Ymin+(Ymax-Ymin)/2
1330
1340
1350
 1360
1370
            1380
 1390
 1400
 1410
            MOVE Xmin-.05*(Xmax-Xmin),Ymin
LABEL USING "K":Ymin
MOVE Xmin-.05*(Xmax-Xmin),Ymax
LABEL USING "K":Ymax
 1420
 1430
 1440
 1450
            CSIZE 5*(Vxmax-Vxmin)/131,.6

MOVE Xmin, Ymin-.05*(Ymax-Ymin)

LABEL USING "K":Xmin

MOVE Xmax, Ymin-.05*(Ymax-Ymin)

LABEL USING "K":Xmax
 1460
 1470
 1480
 1490
 1500
 1510
            MOVE Xmin+(Xmax-Xmin)/3.0.Ymin-.3*(Ymax-Ymin)
CSIZE 4.5*(Vxmax-Vxmin)/131..6
LABEL USING 1550;Sample$.Run_num.f_ov.f_dc.f_fc
IMAGE 18A." RUN #".K.1X.K."C".1X.K."C".1X.K."C"
VIEHPORT Vxmin,Vxmax.Vymin.Vymax
 1520
1530
1540
 1550
1560
 1570
1580
1590
            CLIP Xmin.Xmax,Ymin,Ymax
PEN 1
MOVE 0.0
             RETURN
  1600
 1610 Start_timer:
1620
                                            TS=FNTimeS(TIMEDATE)
                                            CLIP OFF
  1630
             CSIZE 4.5 ~ (Vxmax - Vxmin) / 131..6
             MOVE Xmax.Ymin-.3*(Ymax-Ymin)
LABEL USING "9A.1X.8A":TS.Date$
MOVE 0.5
CLIP Xmin.Xmax.Ymin.Ymax
 1640
1650
1660
  1670
              Init_time=TIMEDATE
```

```
1690
                                                               !PRINT "TIMER INITIALIZED."
 1700
 1710 Read_ad:
                      Rd_start=TIMEDATE-Init_time
                      1720
1730
 1740
 1750
1760
1770
1780
                      R(I)=R(I)+32768
IF R(I)>2047 THEN R(I)=R(I)-4096
R(I)=.005+R(I)
 1790
                      Sum=Sum+R(I)
                      Sum_sq=Sum_sq+R(I)*R(I)
WAIT Wt_time !NOTE WAITS ROUNDED TO NEAREST THOUSANDTH.
 1800
 1810
                      NEXT I
Rav(J)=Sum/N
 1820
 1830
 1840
                      Sq_sum=Sum≠Sum/N
 1850
                      S_sq=(Sum_sq-Sq_sum)/(N-1)
IF S_sq<=0. THEN
 1860
 1870
                      Std_dev(J)=0.
 1380
                      ELSE
                      Std_dev(J)=SOR(S_sq)
END_IF
 1890
 1900
 1910
                      Rd_stop=TIMEDATE-Init_time
 1920
                      Seconds(J)=(Rd_stop+Rd_start)/2.
1930
                      RETURN
                          PLDT Seconds(J).Rav(J)
DISP USING 1960;J.Rav(J).Std_dev(J).Seconds(J)
IMAGE "POINT #".4D.5X."I= ".2D.3D.,5X."STD_DEV= ".2D.3D.5X."=
1940 Plot_inten: 1950
1960
! ".4D.1D." SECS."
1970
                                              RETURN
1980 Knob_thing:
                        ON KEY 1 LABEL "DATA PRINT" GOSUB Data_print
1990
                                                      DISP "END OF EXPERIMENT"
2000
2010
                                                      Tone=81.38
FOR M=1 TO 5
 2020
                                                      BEEF Tone..2
2030
 2040
                                                       Tone=Tone+2
2050
                                                      NEXT M
2060
                                                      MATT
                                                 BEEP
DISP "USE THE KNOB TO MOVE CURSOR THRU THE I
2070
2080
       AND KEY 3 TO PRINT VALUES."
ATA.
2090
                                                 MOVE 0.0
                                                 ON KEY 3 LABEL "CURSOR PRINT" GOTO Cursor_p:
2100
2110
nt
2110
2120
2130
2140
2150
2160 Spin:
2170
2180
                                                 ON KEY 0 LABEL "DISC STORE" GOSUB Disc_store
ON KEY 4 LABEL "QUIT" GOTO Quit
                                                 CLIP OFF
                                                 ON KEY 5 LABEL "PRINT SPOT" GOSUB Print_spot
ON ENDR .! GOSUB Move_birp
GOTO Spin
                                                         RETURN
RETURN
2180
2190 Off_spot:
2200
2210
2220
2230
2240
2250
                       IF Old_spot<>0 THEM
                         DRAW Seconds(Old_spot).Sav(Old_spot)+Space
                       Old_spot=0
END IF
RETURN
```

```
2260 Print_spot:
2270
2280
2290
2300
                                ! Print point number on graph.
LINE TYPE 1
LDIR 90
LORG 2
                                PEN 1
                                IF Old_spot<>0 THEM
MOVE Seconds(Old_spot).Rav(Old_spot)+4*Space
LABEL USING 2340;Seconds(Old_spot)
IMAGE 4D.2D
2310
2320
2330
2340
2350
                                END IF
                                Old_spot=0
PEN 0
2360
2370
2380
2390
                                RETURN
2400 Move_blip:
2410
2420
2430
                                                                              Spot=Spot+KNOBX
                                                                              Spot=INT(Spot)
                                                                            IF Spot-Im (Spot-)
IF Spot>Im_test THEN Spot=Im_test
DISP USING 1960;Spot,Rav(Spot).Stc_
2440
2450
dev(Spot),Seconds(Spot)
2460
2470
                                                                            Space=(Ymax-Ymin)/40
PEN 0
2480
                                                                            GOSUB Off_spot
2490
                                                                            MOVE Seconds(Spot).Rav(Spot)+Space
DRAW Seconds(Spot).Rav(Spot)+3*Spac
2500
2510
                                                                            Old_spot=Spot
                                                                   RETURN
2530 Cursor_prnt:
                                      PRINTER IS 3
2540
2550
                       PRINT USING 1960; Spot.Rav(Spot).Std_dev(Spot).Seconds(Spot)
                                                                                       GOTO Spin
RETURN
                                 OUTPUT 2 USING "#.B";255.75
GCLEAR
RETURN
2570 Clear_screen:
2580
2590
2600 Data_print:
                                !PRINTER IS 9
PRINT "
2610
PROGRAM"
                                                                             GENERAL DATA OUTPUT FROM AD_RLTIME
2620
2630
                                   PRINT USING 2640:Oprs.Dates.Ts
IMAGE "OPERATOR: ".184.9X." DATE: ".8A.9X." TIME: ".8A
2640
                                 PRINT USING 2660:Samples.Run_num
IMAGE "SAMPLE: ".18A.5X." RUN NUMBER: ".K
PRINT USING 2680:T_ov.T_dc.T_fc
IMAGE "OVEN TEMP: ".K.5X."F CELL TEMP:
2650
2660
2670
2680
".K
                                 PRINT USING 2700:Gas_f$.Gas_g$
IMAGE "FLOWMETER-F GAS: ".18A.5X."FLOWMETER-G GAS: ".18A
PRINT USING 2720:Flow_f.Flow_g
IMAGE "STARTING FLOWRATE-F: ".K."CM STARTING FLOWRA
2690
2700
2710
2710
2720
: ".K."CM"
                                                                                                       STARTING FLOWRATE-G
2730 PRINT USING 2740:N.Im_cnst.Test_length
2740 IMAGE 4X.K." SAMPLES AVERAGED OVER(APPROX) ".K." SEC. FOR(AF
2750) ".K." TOTAL SECONDS."
2750
2760
"C"
2770
2780
                                  PRINT USING 2760:Samples.Run_num.T_ov.f_dc.T_fc
IMAGE "GRAPH LABEL: ".18A." KÜN #".E.1X.E."C".1X.K."C",1X.K.
                                  PRINT USING "40A":"COMMENTS: ".CommentS
                                  PRINT
```

SALLES CONTRACTORS AND CONTRACT CONTRACT PROPERTY PROPERTY CONTRACTOR CONTRACTOR CONTRACTOR

```
2790
                                                                                                                                                                                                                                   PRINT "
                                                                                                                                                                                                                                                                                                                                                                                                                               EXCITATION CONDITIONS"
                                                                                                                                                                                                                                  PRINT PRINT USING 2820:Sources
IMAGE "SOURCE: ".18A
PRINT USING 2840:Lam_ex.Bndwth.Dyes
IMAGE "WAVELENGTH: ".K."NM ".10X."BANDWIDTH: ".K."NM",10X."E
                                                                                                                           2800
2810
2820
                                                                                                                             2830
                                                                                                                           2840
YE: "
                                                                                                                                                                                                                                    PRINT USING 2860:P_power,Rep_rate.Power IMAGE K," MJ/PULSE AT ",K,"HZ IS ",K," WATTS."
                                                                                                                            2850
                                                                                                                             2860
                                                                                                                            2870
2880
                                                                                                                                                                                                                                    PRINT
                                                                                                                                                                                                                                   PRINT
                                                                                                                                                                                                                                                                                                                                                                                                                                         EMISSION CONDITIONS"
                                                                                                                             2890
                                                                                                                                                                                                                                    PRINT
                                                                                                                                                                                                                                PRINT USING 2910:Mono$
IMAGE "MONOCHROMATOR TYPE: ",18A
!WAIT 15
                                                                                                                             2900
                                                                                                                          2910
2910
2920
2930
2940
5: ".K."MM"
                                                                                                                                                                                                                                   PRINT USING 2940:Lam_em.Band.Slits
IMAGE "WAVELENGTH: ".K."NM",5X."BANDWIDTH: ".K."NM",5X,"SLIT
                                                                                                                            2950
                                                                                                                                                                                                                                    PRINT
                                                                                                                            2960
2970
2980
                                                                                                                                                                                                                                    PRINT "
                                                                                                                                                                                                                                                                                                                                                                                                                                      DETECTION ELECTRONICS"
                                                                                                                                                                                                                                    PRINT
                                                                                                                                                                                                                                   PRINT USING 2990:Pmts.Hvs.Hvr
IMAGE "PMT: ".10A.10X."SAMPLE-HV: ",K.10X, "REFERENCE-HV: ",
PRINT USING 3010:Proc_insts
IMAGE "PROCESSING INSTRUMENT: ".18A
                                                                                                                             2990
                                                                                                                             3000
                                                                                                                             3010
                                                                                                                           3020 PRINT USING 3030:Gate_time.Ap_dly.Tcs.Tcr
3030 IMAGE "GATE TIME: ",K."MSEC.AP.DELAY: ",K."MSEC.SMPL TIME CN.
ST: ",K."SEC.REF TIME CNST: ",K,"SEC"
3040 PRINT USING 3050:Sens_s,Sens_r
3050 IMAGE "SENSITIVITY OR GAIN - SAMPLE: ",K.SX." REFERENCE: ".
ST: ".K."SEC.REF IIMÉ CMSI: ".K. "SEC"

3040
PRINT USING 3050:Sens_s.Sens_r
1MAGE "SENSITIVITY JR GAIN - SAMPLE: ".K
3050
PRINT USING 3050:Sens_s.Sens_r
1MAGE "SENSITIVITY JR GAIN - SAMPLE: ".K
3080
PRINT USING 3100:J
3100
PRINT USING 3100:J
3100
PRINT USING 3100:J
3110
PRINTER IS 1
PRINTER IS 9
PRINTER IS 9
PRINTER IS 9
PRINTER IS 100:UMB LABEL".Dsc_lbis
3120
RETURN
3130 Disc_store: ! STORE DATA ON DISC
3140
INPUT "ENTER DISC VOLUME LABEL".Dsc_lbis
3150
INPUT "ENTER DISC VOLUME LABEL".Dsc_lbis
3150
PRINTER IS 100:UMB POINT APO EXP FILE:
3160
CREATE BDAT File_names.1651.8
3170
ASSIGN #Path TD File_names.1751.8
3180
OUTPUT #Path:Seconds(*).Rav(*).Std_dev(*)
3190
OUTPUT #Path:Run_nun. N. Test.Cor.Std.
3190
OUTPUT #Path:Run_nun. N. Test.Cor.Std.
3200
PRINTER STORED ON DISC LABELED ".Dsc_
9.Ts.Sample$.Gas_f$.Gas_g$.Comments.Source$.Dye$.Mono$.Pat$
3210
ASSIGN #Path TO *
3220
PRINTER IS 3
440
PRINTER IS 3
450
PRI
                                                                                                                           RETURN
3080 Flow_change: !PRINTER IS 1
3090 PRINT USING 3100: J
3100 IMAGE "FLOW CHANGE AT POINT # ".4D! CAN'T PRINT SECONDS(J)-
GET 0.000 MOST KEY HITS-J OK THOUGH ?????
3110 !PRINTER IS 9
                                                                                                                         RETURN

3130 Disc_store: ! STORE DATA ON DISC

3140 INPUT "ENTER DISC VOLUME LABEL".Dsc_ltls

3150 INPUT "ENTER FILENAME(INCHS).EXAMPLE:. '001_TM".File_name$ !

RTL NOTE -TM OR -PT FOR TIME OR POINT A/D EXP FILES

3160 CREATE BDAT File_name$:.21651.8

3170 ASSIGN @Path TO File_name$:FORMAT OFF

3180 OUTPUT @Path:Seconds(*).Rav(*).Std_dev(*)

3190 OUTPUT @Path:Run_num.N.Test_length.Tm_test.T_ov.T_dc.T_fc,Flc

u_f,Flow_g.Tm_cnst.Lam_ex.Bndwth.Power.Rep_rate.Lam_em.Band.Slits.Hvs.Hcr

3200 OUTPUT @Path:Gate_time.Ap_dly.Tcs.Tcr.Sens_s.Sens_r.Opr$.Date

3.T$.Sample$.Gas_f$.Gas_g$.Comment$.Source$.Dye$.Mono$.Pmt$

3210 ASSIGN @Path TO *

PRINT "DATA STORED ON DISC LABELED ".Dsc_lbls." IN FILE ".File_name$
```

```
END ! END FOR PROGRAM!!!!!!!
DEF FNTime$(Now) !GIVEN 'SECONDS' RETURN 'HH:MM:SS'
                                                            How=INT(Now) MOD 86400
H=Now DIV 3600
M=Now MOD 3600 DIV 60
                                                            S=Now MOD 60
OUTPUT TS USING "#,ZZ.K":H.":".M,":",S
RETURN TS
FNEND
                                                            DEF FNTime(TS) !GIVEN 'HH:MM:SS' RETURN 'SECONDS'
                                                            ON ERROR GOTO Err
ENTER TS:H.M.S
RETURN (3600*H+60*M+S) MOD 86400
OFF ERROR
RETURN TIMEDATE MOD 86400
                                                                    FNEND
```

GENERAL DATA OUTPUT FROM AD_RETIME PROGRAM

OPERATOR: R.LODA DATE: 4/6/83 TIME: 13:53:51
SAMPLE: ANTHRACENE-N2 RUN MUMBER: 1
OVEN TEMP: 159 D CELL TEMP: 155 F CELL TEMP: 164
FLOWMETER-F GAS: NITROGEN FLOWMETER-G GAS: LAB AIR
STARTING FLOWRATE-F: 1CM (40-7/2)STARTING FLOWRATE-G: 0CM
20 SAMPLES AVERAGED OVER(APPROX) 1 SEC. FOR (APPROX) 1000 TOTAL SECONDS.
GRAPH LABEL: ANTHRACENE-N2 RUN #1 155/M 15.00 TOTAL SECONDS. COMMENTS: 75MM LENS-EM-0160 IRIS-EX

EXCITATION CONDITIONS

SOURCE: EXCIMER LASER
NAVELENGTH: JOSHM BANDWIDTH: HIM
10.4166566667 MJ/PULSE AT 24HZ IS HATTS.
58:33 DYE: NONE

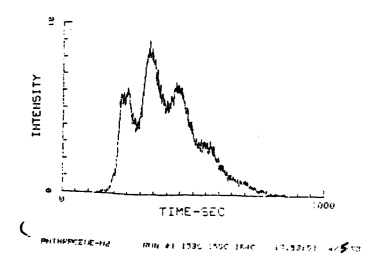
EMISSION CONDITIONS

MONOCHROMATOR TYPE: SPEX .85M 4.233.mm HAVELENGTH: 390MM BANDWIDTH: 9HH SLITS: 1MM

(

DETECTION ELECTRONICS

PMT: RCA C31034 SAMPLE-HV: -1650 REFERENCE-HV: -850 PROCESSING INSTRUMENT: DGA-1 GATE TIME: .1MSEC.AP.DELAY: OMSEC.SMPL TIME CNST: ISEC.REF TIME CNST: .3SEC SENSITIVITY OR GAIN - SAMPLE: 30 REFERENCE: 10



```
.520
020-<del>500</del>nm at 2a/sec
FLOW CHANGE AT POINT #
35000 CHANGE AT POINT #
                              100
                              150
                              200
3600A
FLOW CHANGE AT POINT #
                              250
3700A
FLOW CHANGE AT POINT #
                              300
2800A
FLOW CHANGE AT POINT ≠
3900A
                              350
FLOW CHANGE AT POINT #
                              400
4000A
FLOW CHANGE AT POINT #
4100A
FLOW CHANGE AT POINT # 4200A FLOW CHANGE AT POINT #
                              500
                              550
4300A
FLON CHANGE AT POINT #
4400A
FLOW CHANGE AT POINT #
                              651
4500A
FLOW CHANGE AT POINT =
                              701
4600A
FLOW CHANGE AT POINT #
                              751
4700A
FLOW CHANGE AT POINT #
180
      CHANGE AT POINT #
                              351
4900A
FLOW CHANGE AT POINT = 5000A
```

4/5/83 Run #1

Here, the points are not flow changes, but points at which the monochromater reading was the listed wavelength Value.

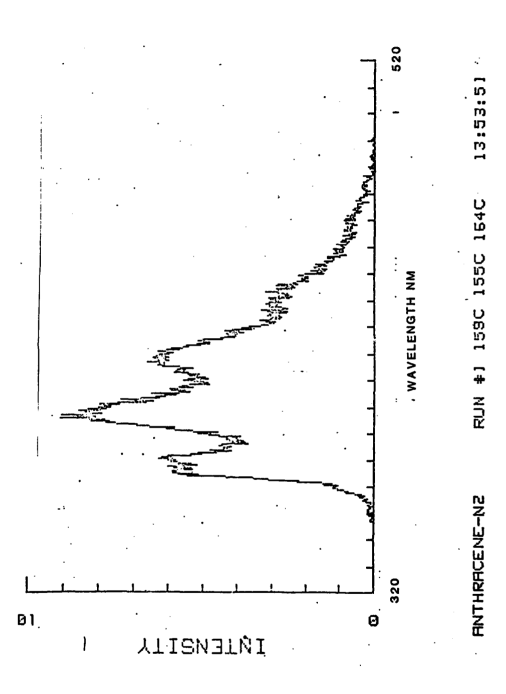
```
100.9 SECS.
151.1 SECS.
201.1 SECS.
                                                                                                                                        .032
.040
.121
                                                                                                   STD DEV-
POINT = 100
POINT = 150
                                                                      .064
                                                                                                                                                                                    201.1
201.0
201.0
350.9
400.8
450.3
500.7
500.5
600.5
POINT # 200

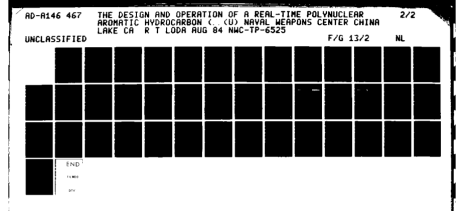
POINT # 250

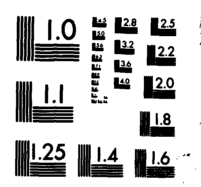
POINT # 350

POINT # 400

POINT # 400
                                                                   1.121
                                                                                                                                                                       AT
                                                                                                                                         .148
.167
.159
                                                                                                                                                                       AT
                                                                  5.902
                                                                  4.811
7.835
5.560
                                                                                                                                                                       AT
                                                                                                                                                                       AT
                                                                                                                                          .094
                                                                                                                                                                       AT
AT
 POINT #
                          450
                                                                   5.993
POINT # 450
POINT # 550
POINT # 560
POINT # 551
POINT # 751
POINT # 751
POINT # 801
                                                                   3.430
                                                                                                                                                                       AT
                                                                   ī.588
949
                                                                                                                                          .048
                                                                                                                                                                       AT
                                                                                                                                         .052
.037
.023
.036
                                                                        .586
                                                                                                                                                                       AT
                                                                      .274
                                                                                                                                                                       AT
                                                                                                                                                                                     751.4
                                                                                                                                                                                    300.3
801.3
851.4
                                                                                                                                                                       ΑŤ
                   # 301
# 851
                                                                       .176
 THIOC
                                                                                                                                           .023
                                                                                                                                                                                     900.3
```







MICROCOPY RESOLUTION TEST CHART NATIONAL BUREAU OF STANDARDS-1963-A

GENERAL DATA OUTPUT FROM AD_RLTIME PROGRAM

DPERATOR: R.LODA DATE: 4/5/83 TIME: 15:18:43
SAMPLE: ANTHRACENE-N2 RUN NUMBER: 2

DVEN TEMP: 159 D CELL TEMP: 155 F CELL TEMP: 164
FLOHMETER-F GAS: NITROGEN FLOHMETER-G GAS: LAB AIR
STARTING FLOHMATE-F: 1CM STARTING FLOHMATE-G: 0CM
20 SAMPLES AVERAGED OVER(APPROX) 1 FLOHMATE-G: 0CM
COMMENTS: RUN #2 159C 155C 164C
COMMENTS: 75MM LENS-EM-0160 IRIS-EX

EXCITATION CONDITIONS

SOURCE: EXCIMER LASER HAVELENGTH: 308NM BANDHIDTH: 1NM 58.333333333 MJ/PULSE AT 24HZ IS 1.4 HATTS.

DYE: NONE

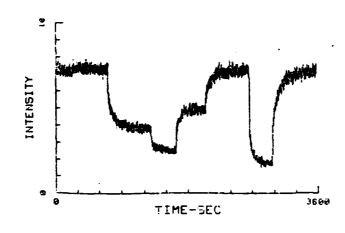
EMISSION CONDITIONS

MONOCHROMATOR TYPE: SPEX .85M

HAVELENGTH: 390NM

DETECTION ELECTRONICS

PMT: RCA C31034 SAMPLE-HV: -1650 REFERENCE-HV: -850
PRZ TSSING INSTRUMENT: DGA-1
GAL TIME: .1MSEC.AP.DELAY: GMSEC.SMP! TIME CNST: 1SEC.REF TIME CNST: .3SEC
SENSITIVITY DR GAIN - SAMPLE: 30 REFERENCE: 10



FLOW AT ICM=40ML/MIN FLOW CHANGE AT POINT # 720 TO 3.6CM-80ML/MIN FLOW TO 3.6CM-8UML/MIN FLOW CHANGE AT POINT # 1302 FLO TO 6.2CM-120ML/MIN FLOW CHANGE AT POINT # 1642 FLOW CHANGE AT POINT # 2051 FLOW CHANGE AT POINT # 2051 FLOW CHANGE AT POINT # 2667 FLOW CHANGE AT POINT # 2667 FLOW TO 8.8CM-160ML/MIN FLOW TO 1.0CM-40ML/MIN FLOW TO 1.0CM-40ML/MIN FLOW TO 1.0CM=40ML/MIN

RUN 42 1590 1550 1840

GENERAL DATA OUTPUT FROM AD_RLTIME PROGRAM

OPERATOR: R.LODA DATE: 4/5/83 TIME: 16:30:22
SAMPI E: ANTHRACENE-N2-AIR RUN NUMBER: 3
OVA TEMP: 159 D CELL TEMP: 155 F CELL TEMP: 164
FLOWMETER-F GAS: NITROGEN FLOWMETER-G GAS: LAB AIR
STARTING FLOWRATE-F: 1CM STARTING FLOWRATE-G: 0CM
20 SAMPLES AVERAGED OVER(APPROX) 1 SEC. FOR (APPROX) 3600 TOTAL SECUNDS.
GRAPH LABEL: ANTHRACENE-N2-AIR RUN #3 1590C 1550C 164C
COMMENTS: 75MM : ENS-EM-0160 IRIS-EV COMMENTS: 75MM LENS-EM-0160 IRIS-EX

EXCITATION CONDITIONS

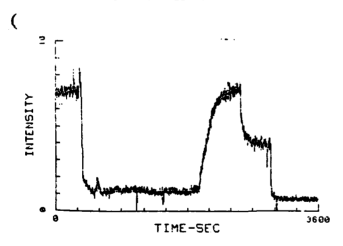
SOURCE: EXCIMER LASER
HAVELENGTH: 308NM BANDHIDTH: 1NM
58.33333333333 MJ/PULSE AT 24HZ IS 1.4 HATTS. DYE: NONE

EMISSION CONDITIONS

MONDCHROMATOR TYPE: SPEX.85M OFFICE SHAPELENGTH: 390NM BANDHIDTH: 9HM SLITS: !MM

DETECTION ELECTRONICS

PMT: RCA C31034 SAMPLE-HV: -1650 REFERENCE-HV: -850 PROCESSING INSTRUMENT: DGA-1 GATE TIME: .1MSEC.AP.DELAY: OMSEC.SMPL TIME CNST: 1SEC.REF TIME CNST: .3SEC SENSITIVITY OR GAIN - SAMPLE: 30 REFERENCE: 10



FINTHRRCENE-NZ-RIR FUN #3 1590 1550 1640

SECRETAL PROPERTY SECRETAL CONTRACT CONTRACT SECRE

```
FEON TO 1.0CH=40ML/MIN
FLOW CHANGE AT POINT # 238
            N2-0.AIR-1.3CM-40ML/MIN
NOTE TIME DELAY TO PT 420 OR SO FOR AIR GUENCH-TUBING VOLUMEAFLON RATE?
FLOW CHANGE AT POINT # 592
PROBLEMS WITH SETTING FLOW RATE-PRESSURE FLUCTUATIONS
FLOW CHANGE AT POINT # 713
PROBLEMS WITH SETTING FLOW RATE-PRESSURE FLUCTUATIONS
FLOW CHANGE AT POINT # 811
PROBLEMS WITH SETTING FLOW RATE-PRESSURE FLUCTUATIONS FLOW CHANGE AT POINT # 1100
TIL BLOCKING BEAM TO SAMPLE
FLOW CHANGE AT POINT # 1479
FROBLEMS WITH SETTING FLOW RATE-PRESSURE FLUCTUATIONS
FLOW CHANGE AT POINT # 1842
PLGW CHANGE AT POINT # 1842
FLOW N2-ICM-40ML/MIN.AIR-0
TOTE TIME DELAY TO PT 1980 OR SO FOR AIR QUENCH-TUBING VOLUMEAFLOW RATE?
FLOW CHANGE AT POINT # 2543
FLOW N2-3.6CM-80ML/MIN.AIR-0
FLOW CHANGE AT POINT # 2903
FLOW N2-0.AIR-4.4CM-80ML/MIN
TOTE TIME DELAY TO FT 2966 OR SO FOR AIR QUENCH-TUBING VOLUMEAFLOW RATE?
FLOW CHANGE AT POINT # 3083
FLOW CHANGE AT POINT # 3083
FLOW CHANGE AT POINT # 3083
RTL BLOCKING BEAM TO SAMPLE
 LASER POWER AT IN AT END OF EXPERIMENT
                                                                                                                                                                                              4/5/83
                                                                                                                                                  37.6 SECS.

44.5 SECS.

52.6 SECS.

105.5 SECS.

169.3 SECS.

233.0 SECS.

237.0 SECS.

267.0 SECS.

276.9 SECS.

284.9 SECS.

284.9 SECS.

284.8 SECS.
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STO DEY=
STO DEY=
  POINT =
POINT =
                        38
45
63
                                                      5.974
                                                                                                                                                                                              Run # 3
                                                                                                                .163
                                                      6.569
6.939
                                            ] =
] =
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                                                                                  STD DEV=
STD DEV=
STD DEV=
STD DEV=
               #
                                                                                                                                       ΑT
  POINT
20 ( )
                                                                                                               .098
.122
.095
.122
 POINT = 106
POINT = 238
POINT = 238
POINT = 238
POINT = 293
POINT = 308
POINT = 308
POINT = 308
POINT = 352
POINT = 357
POINT = 367
POINT = 369
POINT = 369
POINT = 377
POINT = 388
                                                      5.901
6.920
                                                                                                                                       AT
AT
                                                                                                                                       AT
AT
                                                       7.074
                                                      6.956
                                                                                 STD DEV-
STD DEV-
                                                      6.995
7.152
6.647
                                                                                                                .104
                                                                                                                .106
                                                                                                                                       аT
                                                                                                                                       AT
AT
AT
                                                                                 STD DEV-
                                                                                                               .054
                                                       6.827
                                                       6.314
7.074
                                                                                                                ,049
                                                                                                                                                    300.8
                                                                                                               .056
.161
.072
                                                                                                                                                  306.7 SECS.
320.7 SECS.
330.6 SECS.
                                                                                                                                        AT
                                                                                                                                       AT
AT
                                                      6.918
7.855
8.235
7.365
7.506
6.216
3.520
2.859
2.013
1.828
                                                                                                                .086
                                                                                                                                       AT
AT
                                                                                                                                                   335.6
342.5
351.5
                                                                                                                . 134
                                                                                                                                                   367.4
367.4
370.4
                                                                                                                .146
                                                                                                                                        AT
                                                                                                                .071
                                                                                                                                       AT
AT
                                                                                                                                                   375.4 SECS.
381.3 SECS.
                                                                                                                .058
                                                                                                                                        AT
                                                                                                                                                                SECO.
                                                                                                                                                  388.3
401.2
408.2
417.1
                                                                                                                .036
                                                                                                                .091
.123
.038
 POINT
                # 403
                                                       1.701
                                                                                                                                        ĄΤ
                                                       1.708
1.627
                                                                                                                                       AT
  29INT # 410
 POINT # 419
                                                                                                                                        ΑŢ
                                                                                                               125. -
                                                                                                                                                   430.1
                                                                                                                                                                  SECS
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STD DEV-
STD DEV-
STD DEV-
                                                                                                      607.0 SECS.
658.5 SECS.
POINT ≠ 608
                                      1.301
1.227
.994
                                                                              .040
20 K . .
               559
591
732
                                                                                                                 SECS.
                                                                              .080
                                                                                                      690.4
731.4
784.1
                                                                                                                 SECS.
SECS.
SECS.
                                                         STD DEV-
                                                                               .040
                                                                                              AT
           =
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- INIO
                              I-
                                        .963
                                                                              .931
                                      1.241
1.306
1.367
1.435
POINT # 785
                                                         STD DEV-
                                                                              .030
                                                                                               AT
                                                         STD DEV-
                                                                                                                 SECS.
SECS.
                              1=
POINT # 842
                                                                              .077
                                                                                               ΑT
                                                                                                      841.0
                                                                                                    865.9
1043.2
                                                                              .054
POINT #
               367
                                                        STD DEV-
20INT =1045
                                                                                               ΑT
                               <u>[</u> =
                                      1.344
                                                                              .093
                                                                                                    1193.1
                                                                                                                 SECS.
 201NT ±1195
                                                                                               AT.
                               Ī.
                                                                                                    1284.6
1361.2
1436.8
1508.7
20INT #1287
                                                         STD DEV-
                                                                                                                 SECS
SECS
                                                                                               AT
                                                                              .054
20INT =1364
                                                                              .083
                                                                DEV-
                                                                                               AT
                               I=
                                      1.140
                                                         STD
                                      1.105
1.137
1.234
1.099
                                                         STD
                                                                                                                 SECS.
SECS.
POINT #1440
                                                                DEV-
 POINT #1512
                                                                DEY-
                                                                               .058
                                                         STD DEV=
STD DEV=
STD DEV=
STD DEV=
                                                                              .ŭ46
POINT #1582
                                                                                                    1578.3
                                                                                               AT
                                                                                                    1670.8
1830.4
1835.4
                                                                                                                 SECS.
SECS.
POINT #1575
                                                                              .052
                                                                                               AT
                               Į=
POINT #1835
                               l=
                                      1.300
                                                                              .043
                                                                                               AT
 POINT #1840
                                      1.176
                                                                              .034
                                                                                               ΑT
                                                         STD
                                                                DEV=
                                                                              .090
                                                                                                     1837.4
                                                                                                                 SECS.
 POINT #1842
                                                                                               AT
 20INT ≠1886
                                      1.436
                                                                                               AT
                                                         STD DEV-
                                      1.319
                                                                                                                 SECS.
 20INT =1895
                                                                              .058
                                                                                                     1890.6
                                                                                               AT
                                                                                                    1900.6
 20INT =1305
                                                                              .038
                                      1.443
                                                                                               ΑŢ
                                                                                                                 SECS.
 POINT ≠1921
                                      1.260
                                                                              .046
 ≎ดีมีที่ ∍เลริธ์
                                                                              .056
.028
                                                                                                     1931.6
                                       1,421
                                                                                               AT
 POINT #1951
                                      1.352
1.300
2.341
2.580
2.583
                                                                                                                 SECS.
SECS.
                                                                                                    1946.5
                                                                                               ĤΤ
                                                                                                    1960.4
980.3
 20INT ≠1965
                                                                               .046
                                                                                               AT
 20INT #1985
                                                                              645
                                                                                               AT.
POINT #1995
POINT #1996
PO( #2011
POINT #2026
POINT #2143
POINT #2281
                                                                                                                 SECS.
SECS.
SECS.
SECS.
                                                                               .095
                                                                                                     1991.5
                                                                                               AT
                                                                                                    1991.5
2006.4
2021.3
2053.2
2137.7
2274.9
                                                                              .041
                                                                                                                                     4/5/83
                                                                                               AT
                                      3.309
                                                         STU DEV-
                                      3.584
5.504
                                                         STD DEV=
STD DEV=
STD DEV=
                                                                              .979
                                                                                               AT
                                                                              .091
                                      6.845
                                                                                               AT
POINT #2388
POINT #2470
POINT #2527
POPT #2568
POUT #2624
                                                                                                    2381.3 SECS.
2462.9 SECS.
2519.7 SECS.
                                      7,120
                                                         STD DEV-
                                                                               .077
                                                                                               ĤΤ
                                      7.103
7.221
5.146
4.324
3.948
                                                         STD DEV-
                                                                              .091
                                                                                                   2519.7 SECS.
2560.8 SECS.
2616.5 SECS.
2696.3 SECS.
2774.8 SECS.
2837.5 SECS.
2876.3 SECS.
2893.2 SECS.
2894.3 SECS.
2939.1 SECS.
2965.5 SECS.
2979.5 SECS.
                                                                              .063
                                                                                               AT
                                                         STD DEV-
STD DEV-
STD DEV-
                               Ţ.
                                                                              .068
                                                                                               AT
POL #2624
POINT #2704
                               Ī=
                                                                              .084
                                                                                               AT
                                                                              .092
.044
.039
                                                                                               AT
                                                         STD DEV-
STD DEV-
STD DEV-
POINT #2783
                                      4.041
                                                                                               AT
 POINT #2846
                                      3.639
3.773
                                                                                               AT
POINT #2885
                                                                                               AT
POINT #2902
POINT #2903
POINT #2948
                                                         STD
                               I=
                                      4.017
                                                                DEV=
                                                                              .111
                                                                                               AT
                                      3.766
4.176
                                                                DEV-
                                                                              .113
                                                                              .123
                                                         STD
                                                                DEV-
                                                                                               AT
                                      1.093
.961
.773
                                                         STD DEV-
STD DEV-
STD DEV-
 PDINT #2974
                                                                                               АT
POINT #2988
POINT #3017
                                                                                               ΑT
                                                                DEV-
                                                                              .038
                                                                                               AT
                                                                                                     5.8906
                                                                                                    3034.5 SECS.
3034.5 SECS.
3061.4 SECS.
3098.2 SECS.
3131.0 SECS.
                                         .625
.768
.739
                                                         STD DEV-
STD DEV-
STD DEV-
 POINT #3043
                                                                              .051
                                                                              .049
.023
.034
.045
.022
.016
 20INT ≠3070
 POINT #3107
                                                                                               AT
                                        .656
.613
.324
                                                         STD
 POINT #3140
                                                                DEV=
                                                                                               AT
                                                                                                    3131.0
3161.6
3374.7
3588.5
2567.6
3588.5
 POINT #3171
                                                                DEV-
                                                                                                                 SECS.
SECS.
SECS.
                                                                                               AT
 POINT #3385
                                                         STD
                                                                DEV-
                                                                                               AT
POINT #3500
POINT #3579
                                        .768
.769
.768
                                                         STD
STD
                                                                DEV=
                                                                                               AT
AT
POINT #3600
                                                                DEV-
                                                                               .618
```

NWC TF 6525

Appendix B

SPEX DATAMATE SPECIFICATIONS

FLUOROLOG°2

SPECIFICATIONS

DATA PROCESSING AND STORAGE - SPEX DATAMATE

The FLUOROLOG comes standard with the DATAMATE (DM1) scan controller/data processor equipped with options. DM105-data storage/graphics/processing/radiometric correction; DM102-PC acquisition module. DM103-DC acquisition module, 4. DM101-input channels; DM104-second High voltage supply, and 1457B-coupled y-t recorder or DM112 plotter. For complete details and specifications refer to the DATAMATE brochure.

SELF-DIAGNOSTICS AND OPERATIONAL ERROR CHECK

- A Power on initiates primary hardware check. Error codes are displayed on keyboard LED for indication even in event of CRT failure. B Internal logic test and software test with errors displayed on CRT. C Continuous monitor of all keyboard operations for compatibility with hardware and software demands previously made. Errors displayed in STATUS field.

SPECTROMETER CONTROL

Scans excitation and/or emission spectrometers with 0.02 nm step resolution. Manual control through forward/reverse toggle switches. \sim

stallen masses et estation for the second consistence of the second property of the second property of the second property of the second consistence of the second consistency

- 1 Continuous
 2 Burst 1 dependent on integration time and increment.
 3 Burst 2 requires trigger for move to next increment.
 4 Burst 3 requires trigger to initiate integration.
 5 Burst 4 requires triggers to initiate integration and movement.
 6 Set slews to chosen position and halts.

- B Multiple Group Scanming
- 1 1-4 groups with independent start, stop, integration and increment (or

Four inputs are available; Two chopped photon counting (\$1, \$2); two chopped direct current (R1, R2).

A Mainframe Data Input

Integration times 0.1 to 800 sec.
Integration time accuracy 0.01%
Data word length 24 bit mentises. 8 bits for exponent and sign
Data word max, significant digits 8
Max. counts data point 2: 10 ** (bipolar)
Max Count Pate (DM101) 25 MHz

8 Data Acquisition Modes

Signats can be acquired and combined in real time according to the mod listed below.

\$1 K81 \cdot 81 R1 S1/R1
\$1/R2 K(\$1/R1) \cdot (\$2/R2)

KS1 - S2 S1/R1 K(S1/R1) - (S2/R2) (S1/R1)(R2/S2) \$1 and \$2 \$1/R1 and \$2/R2

DATA STORAGE, GRAPHICS, RADIOMETRIC CORRECTION PROCESSING

A Data Memory

4000 words RAM, 32 bits
Floating-point formatted
Word length, 24-bit mantissa, 8-bit exponent and sign
Precision 1 part in 16,000,000
8 significant digits, maximum
Maximum value 10 exp. 18
Variable-length files, 8 maximum

B Processing

Anthmetic Combinations of Spectra: +, ~, x, =
Radiometric Correction Program
Weighted Smoothing (Savitsky/Golay — 5, 9, 13, 17, 21 points)
Derivatives (first and second: weighted)

Derivatives (first and second: weighted)
Indefinite Integrals
Definite Integrals and Peak Areas
Logal Antilogs
Shifts of Spectral Axes
Point-by-point insertion or Modification of Spectra

C Graphic Display

Annotated display of any 2 files simultaneously one file intensified Real-time display of data acquisition Labeled axes with scales File tabels maximum 40 characters each file Files displayed need not colincide X-axis Spectra can be shifted during display Two independent cursors with coordinates displayed Vertical/Independent cursors with coordinates displayed Vertical/Independent cursors with coordinates displayed Vertical/Independent cursors with coordinates displayed

Vertical/horizontal expand or contract between cursors Output to recorder of data between cursors

D Multitasking

Simultaneous Data Input Oata Processing Data Output

Two spectral curves can be stored and retained on power off for paseine flattening or radiometric correction. Optional celibrated temps are available from SPEX for generation of correction factors.

Celibration curve can be entered point by point, by scan, or by processing, 250 celibration points maximum.

Automatic interpolation between points

Correction can be in real time or on stored data

ACCESSORIES

DM107

A full line of accessories are available for data storage, processing and output

DM110/11 Single/dual disk for data storage

DM106 Data memory expansion to 16000 points

RS232 interface

DM109

CRT video copier DM112/113 Alphanumeric digital plotter 81/2 x 11 or 17 1457B Coupled y-t recorder

Appendix C
DIFFUSION RATE PROGRAM

```
! PROGRAM "DIFF_RATE". R LODA 4/7/93
!MODIFIED FOR ADDITIONAL OUTPUT 2/2/84
!MODIFIED !/31/84 FOR NEW PPM CALC AND MG/CU-M OUTPUT
          ISEE HNDBK ENVIR. DATA ON ORG. CHEM. ED. BY KAPEL VERSCHUEREN IVAN NOSTRAND REINHOLD CO. 1983 PG 41 FOR FORMULAE PPM BY VOLUME IAND MASS IN MG/CU-M (CONC.)
INPUT "ENTER MOLECULAR WEIGHT OF GAS.SOLID(G/MOLE)".M1.M2
INPUT "ENTER HARD SPHERE COLLISION DIAMETER(ANGSTROMS)",D12
          D12=D12=1.E-8
          INPUT "ENTER SOLID TEMPERATURE IN DEGREES C", To
110
          Tk = Tc + 273.1
          P=9.33E+5 !:
Pressure=700
                            !700TORR ATMOSPHERIC PRESSURE AT NWC IN DYNES*CM-2
:20
:30
          A=3/(8*SQR(2*PI)*6.023E+23)
140
 150
          PRINT
          PRINT "
                            OUTPUT FROM DIFF_RATE PROGRAM WITH P ="", Pressure," TORR"
160
170
          PRINT
          PRINT "A= ".A
:80
          B=((8.3144761E+7*Tk) 1.5)/P
FRINT "B= ".B
190
          C=SQR((M1+M2)/(M1+M2))/(D12/2)
PRINT "C= ",C
         P=A+B+C
PRINT "DIFFUSION COEFFICIENT= ".D."CM+2/SEC-MOLECULE"
PRINT "FOR T= ".Tc."C. COLLISION DIAMETER= ".D12+1.E+8."ANGSTROMS"
PRINT "AND M1= ".M1."G/MOLE, M2= ".M2."G/MOLE"
L=11.5 !DIFF LENGTH FOR OUR SYSTEM IN CM
      ?50
360
          Dif_rate=R*R1*1.E-3

PRINT "DIFFUSION RATE~ ".Dif_rate."G/SEC AT ".Vap_pres."TORR OF SOLID"
INPUT "ENTER FLUORESCENCE CELL TEMPERATURE (DEGREES C)".Tfcc
PRINT "FLUORESCENCE CELL TEMPERATURE IS ".Tfcc." DEGREES C"
 390
400
410
          PRINT
          Tfck=Tfcc+273.1
          INPUT "ENTER FLOWRATE OF GAS IN ML/MIN". Gas_flowm
         Gas_flows=Gas_flowm/60
Conc=(Dif_rate/Gas_flows) *1.E+9
PRINT "CONCENTRATION= ".Conc."MG/CU-M AT ".Gas_flowm," ML/MIN"
450
460
470
480
          49û
          Ppm=(760*.0820575*Tfck*Conc)/(700*M2)
          PRINT "THE CONCENTRATION IS ".Conc." MG/CU-M. OR ".Ppm." PPM"
PRINT "THE CONCENTRATION IS ".Conc." MG/CU-M. OR ".Ppm." PPM"
PRINT "FOR THE ABOVE SET OF CONDITIONS"
510
520
          PRINT
560
          GÖTÖ 430
END
```

```
OUTPUT FROM DIFF_LASL PROGRAM WITH P -
                                                                                                                                     585
                                                                                                                                                               TORR
                             2.48386775943E-25
6.05649789236E+9
1.35212433269E+14
C= 1.35212433269E+14

DIFFUSION COEFFICIENT= .203407362808

FOR T= 65.5 C, COLLISION DIAMETER= 3.9 ANGSTROMS

AND M1= 29 G/MOLE, M2= 128 G/MOLE

TIME FOR STEADY STATE= 325.086560719

DIFF. LENGTH= 11.5 CM. CROSS SEC. AREA .126 CM2

DIFFUSION RATE= 3.69665135774E-8 G/SEC AT 2.73 TORR OF SOLID

FILIORESCENCE CELL TEMPERATURE IS 65.5 DEGREES C
 CONCENTRATION= 55.449770366 MG/CU-M AT 40 THIS IS 3.54606758196 MG/CU-M = 1 PPM AT THIS T.P. AND M2 THE CONCENTRATION IS 55.449770366 MG/CU-M, OR 15.6369750673 PPM FOR THE ABOVE SET OF CONDITIONS
                                                                                                                                                                                            ML/MIN
 CONCENTRATION= 18.4832567887 MG/CU-M AT 120 THIS IS 3.54606758196 MG/CU-M = 1 PPM AT THIS T.P., AND M2 THE CONCENTRATION IS 18.4832567887 MG/CU-M, OR 5.21232502243 PPM FOR THE ABOVE SET OF CONDITIONS...
                                                                                                                                                                                            ML/MIN
 CONCENTRATION= 13.8624425915 MG/CU-M AT 160
THIS IS 3.54606758196 MG/CU-M 1 PPM AT THIS T.P. AND M2
THE CONCENTRATION IS 13.8624425915 MG/CU-M. OR 3.90924376682 PPM
                                                                                                                                                                                            ML/MIN
 3.90924376682 PPM
FOR THE ABOVE SET OF CONDITIONS
 CONCENTRATION= 8.21478079497 MG/CU-M AT 270
THIS IS 3.54606758196 MG/CU-M = 1 PPM AT THIS T.P. AND M2
THE CONCENTRATION IS 8.21478079497 MG/CU-M, OR
2.31658389886 PPM
FOR THE ABOVE SET OF CONDITIONS
                                                                                                                                                                                            ML/MIN
 CONCENTRATION= 6.93122129575 MG/CU-M AT 320
THIS IS 3.54606758196 MG/CU-M = 1 PPM AT THIS T.P. AND M2
THE CONCENTRATION IS 6.93122129575 MG/CU-M, UR
1.95462188341 PPM
                                                                                                                                                                                            ML/MIN
  FOR THE ABOVE SET OF CONDITIONS
         OUTPUT FROM DIFF_LASE PROGRAM WITH P -
                                                                                                                                     492
                                                                                                                                                               TORR
                              2.48386775943E-25
7.20131864649E+9
                               1.35212433269E+14
 C* T.35212433263E+14

DIFFUSION COEFFICIENT* .241856145359 CM+2/SEC-MOLECULE
FOR T* 65.5 C. COLLISION DIAMETER* 3.9 ANGSTROMS

AND M1* 29 G/MOLE, M2* 128 G/MOLE

TIME FOR STEADY STATE* 273.406325491 SECONDS

DIFF. LENGTH * 11.5 CM, CROSS SEC. AREA * .126 CM2

DIFFUSION RATE* 4.39735287896E-8 G/SEC AT 2.73 TORR OF SOLID

FLUORESCENCE CELL TEMPERATURE IS 55.5 DEGREES C
                                                                                                                                  CM+2/SEC-MOLECULE
3.9 ANGSTROMS
G/MOLE
  CONCENTRATION= 32.9801465922 MG/CU-M AT 80
THIS IS 2.98233376124 MG/CU-M = 1 PPM AT THIS T.P. AND M2
THE CONCENTRATION IS 32.9801465922 MG/CU-M, OR
                                                                                                                                                                                            ML/MIN
 11.058502915 PPM
FOR THE ABOVE SET OF CONDITIONS
```

TEAM PROTECTION OF THE PROPERTY OF THE PROPERT

OUTPUT FROM DIFF_RATE PROGRAM WITH P =	700	TORR
A= ' 2.48386775943E-25 B= 7.29872649457E+9 C= 7.69517242317E+13 DIFFUSION COEFFICIENT= ,1395063304 FOR T= 159 C, COLLISION DIAMETER= AND M1= 28 G/MOLE, M2= 1 TIME FOR STEADY STATE= 473.9928272 DIFF, LENGTH = 11.5 CM, CROSS S DIFFUSION RATE= 2.27536938515E-8 G/FLUORESCENCE CELL TEMPERATURE IS	185 CM+2/SEC-1 178 G/MOLE 207 SECONDS SEC. AREA = 7SEC AT 2.25 164 DEGREES	10LECULE ANGSTROMS .126 CM2 TORR OF SOLID
CONCENTRATION= 34.1305407772 MG THIS IS 4.57093833206 MG/CU-M = 1 THE CONCENTRATION IS 34.13054077 7.46685653968 PPM FOR THE ABOVE SET OF CONDITIONS	J/CU-M AT PPM A! THIS T.P. 772 MG/CU-M.	40 ME/MIN AND M2 OR
CUNCENTRATION= 17.0652703886 MC THIS IS 4.57093833206 MG/CU-M = 1 THE CONCENTRATION IS 17.06527038 3.73342826984 PPM FOR THE ABOVE SET OF CONDITIONS	I PPM AT THIS T.P. 886 MG/CU-M.	80 ML/MIN AND M2 UR
CONICENTRATION= 11.3768469257 MG THIS IS 4.57093833206 MG/CU-M = 1 THE CONCENTRATION IS 11.37684692 2.48895217989 PPM FOR THE ABOVE SET OF CONDITIONS	GZCU-M AT	120 MEZMIN AND M2 OR
CONCENTRATION= 8.5326351943 MG THIS IS 4.57093833206 MG/CU-M = 1 THE CONCENTRATION IS 8.532635194 1.86671413492 PPM FOR THE ABOVE SET OF CONDITIONS	F/CU-M AF PPM AT THIS T.P. F3 MG/CU-M.	160 MEZMIN AND M2 OR
CONCENTRATION= 6.82610815544 MG THIS IS 4.57093833206 MG/CU-M = 1 THE CONCENTRATION IS 6.826108155 1.49337130794 PPM FOR THE ABOVE SET OF CONDITIONS	G/CU-M A: PPM AT THIS T.P. MG/CU-M.	200 ME/MIY AND M2 OR
CONCENTRATION - 4.55073877029 MC THIS IS 4.57093833206 MG/CU-M = 1 THE CONCENTRATION IS 4.550738770 .995580871958 PPM FOR THE ABOVE SET OF CONDITIONS	G/CU-M AT PPM AT THIS T.P. DE MG/CU-M.	300 ML/MIN AND M2 OR

Air

OUTPUT FROM DIFF_RATE PROGRAM WIT	TH P = 700	TORR
A= 2.48386775943E-25 B= 7.29872649457E+9 C= 7.57966365927E+13	1004.005/I (M40./600)	MOLECULE
DIFFUSION COEFFICIENT : 13741 FOR T= 159 C. COLLISION DIAN AND M1= 29 G/MOLE, M2= TIME FOR STEADY STATE 481.21	1ETER= 5.14 178 G/MOLE 16145817 SECONDS	ANGSTROMS
TIME FOR STEADY STATE 481.21 DIFF. LENGTH - 11.5 CM. CF DIFFUSION RATE 2.24121484115E-9 FLUORESCENCE CELL TEMPERATURE IS	ROSS SEC. AREA = B G/SEC AT 2.25 164 DEGREES	.126 CM2 TORR OF SOLID C
OBNCENTRATION= 33.6182226172 THIS IS 4.57093833206 MG/CU- THE CONCENTRATION IS 33.618 7.35477492257 PPM FOR THE ABOVE SET OF CONDITIONS	-M = 1 PPM AT THIS T.P.	AND M2
CONCENTRATION= 16.8091113086 THIS IS 4.57093833206 MG/CU- THE CONCENTRATION IS 16.809 3.67738746129 PPM FOR THE ABOVE SET OF CONDITIONS	-M = 1 PPM AT THIS T.P. 91113086 MG/CU-M.	. AND M2

```
OUTPUT FROM DIFF_RATE PROGRAM WITH P'=
                                                                                                 700
                                                                                                                    TORR
A= 2.48386775943E-25
B= 5.06292064969E+9
C= 1.35212433269E+14
DIFFUSION COEFFICIENT= .17003809062 CFOR T= 65.5 C. COLLISION DIAMETER= AND M1= 29 G/MOLE, M2= 128 GFOR STEADY STATE= 388.883454049 SFOR STEADY STATE= 388.883454049 SFOR STEADY STATE= 3.08902175695E-8 G/SEC AT FLUORESCENCE CELL TEMPERATURE IS 65.5
                                                                                                CM+2/SEC-MOLECULE
                                                                                                3.9
G/MOLE
                                                                                                                  ANGSTROMS
                                                                                      AREA = .126 CM2
AT 2.73 TORR OF SOLID
                                                                                                  DEGREES C
CONCENTRATION= 23.1676631771 MG/CU-M AT 80 THIS IS 4.24315779038 MG/CU-M = 1 PPM AT THIS T.P. AND M2 THE CONCENTRATION IS 23.1676631771 MG/CU-M, OR 5.46000510037 PPM
                                                                                                                                         ML/MIN
 FOR THE ABOVE SET OF CONDITIONS
        OUTPUT FROM DIFF_SEA PROGRAM WITH P =
                                                                                                 760
                                                                                                                    TORR
CONCENTRATION= 21.3294763784 MG/CU-M AT 80
THIS IS 4.60685702956 MG/CU-M = 1 PPM AT THIS T.P. AND M2
THE CONCENTRATION IS 21.3294763784 MG/CU-M PPM
                                                                                                                                         ML/MIN
FOR THE ABOVE SET OF CONDITIONS
      OUTPUT FROM DIFF LASE PROGRAM WITH P =
                                                                                                 585
                                                                                                                    TORR
                     2.48386775943E-25
6.05649789236E+9
1.35212433269E+14
 B =
C= 1.332124332032
DIFFUSION COEFFICIENT= .203407362808
FOR T= 65.5 C, COLLISION DIAMETER=
AND M1= 29 G/MOLE, M2= 128
                                                                                               CM+2/SEC-MOLECULE
3.9 ANGSTRO
G/MOLE
AND M1= 29 G/MOLE, M2= 128 G/MOLE, M2= 128 TIME FOR STEADY STATE= 325.086560719 SECONDS
DIFF. LENGTH = 11.5 CM, CROSS SEC. AREA = .126 CM2
DIFFUSION RATE= 3.69665135774E-8 G/SEC AT 2.73 TORR OF SOLID
FUNDESCENCE CELL TEMPERATURE IS 65.5 DEGREES C
 THIS IS 3.54606758196 MG/CU-M = 1 PPM AT THIS T.P. AND M2
THE CONCENTRATION IS 27.724885183 MG/CU-M, DR
7.81848753864 PPM
                                                                                                                                         ML/MIN
 FOR THE ABOVE SET OF CONDITIONS
```

Appendix D

SPEX DIFFUSION RATE PROGRAM

```
M1 GAS G/MOLE
1) 29 TO RO
2) 128 TO R1
3) 3.9 TO R2
                                                           M2 SOLID G/MOLE
                                                           HARD SPHERE DIA-A
                                                           ATM FRESSURE-DYNES/CM2
4) 7.7994E+05 TO R3
                                LASL (585 TORR)
                                                           TEMPERATURE-K
5) 338.6 TO R4
                                                           DIFF CELL AFSA-CM<sup>2</sup>
DIFF TUBE LENGTH-CM
VAP PRESSURE-TORR
6) .126 TO RS
7) 11.5 TO R6
8) 2.73 TO R7
9) 80 TO R8
                                                           FLOW RATE-ML MIN
10) 2 # 3.14159 TO R9
11) SOT R9 TO R9
11) SUL RY 10 RY
12) 8 * R9 TO R9
13) & 023 * R9 TO R9
14) 3 / R9 TO R9
15) 8.315E07 * R4 TO RA
                                                           R IN (DYNE-DM) / (MOLE-K)
16) SOT RA TO RB
17) RA * RB TO RB
18) RB /RI TO RA
19) RO # R1 TO RB
20) R0 + R1 TO RC
21) RC / RP TO RB
22) SGT RE TO RB
25) R2 * R2 TG RC
 24) RB / RC TO RB
25) R9 # RA TO RC
26) RC * RB TO RC
27) 1E-07 * RC TO RC
                                                           DIFF COEF-CM2 (SEC-MOLECULE)
28) R6 * R6 TO RD
29) RD / 2 TO RD
                                                           STEADY STATE TIME-SEC
30) RD / RC TO RD
31) RC # R5 TO RE
32) RE # R1 TO RE
 33) 585 * RE TO RE
34) RE / 760 TO RE
35) RE / .08205 TO RE
36) RE / R4 TO RE
37) RE / R6 TO RE
 38) 585 - 87 TO RF
39) 1 / RF TO RF
 40) 585 # RF TO RF
41) LOG RF TO RF
42) 2.303 * RF TO RF
```

$$W(q)*10^{3} (mg/g) = RT$$

$$ppm = \frac{1}{V(1)*10^{10} (m^{3}/1)} RM_{10}^{2}$$

$$PM_{10}^{2}$$
(D.1)

NWC TP 6525

CON	ZERSION FA			492		TORR
	1PPM	= X MG/CL	J-M FROM	TABLE		
T(C)	CIOHB	C11H10	C14H10	C16H10	C18H12	C20H12
20.000	3.447	3.824	4.793	5.439	6.140	6.786
25.000	3.389	3.760	4.713	5.348	6.037	6.672
30.000	3.333	3.698	4.435	5.260	5.937	6.562
35.000	3.279	3.638	4.560	5.175	5.841	6.455
40.000	3.227	3.579	4.487	5.092	5.747	6.352
45.000	3.176	3 .5 23	4.416	5.012	5.657	6.252
50.000	3.127	3.469	4.348	4.934	5.569	6.156
55.000	3.07 9	3.416	4.282	4.859	5.484	6.062
60.000	3.033	3.364	4.217	4.786	5.402	5.971
65. 000	2.988	3.315	4.155	4.715	5.322	5.882
70.000	2.944	3.266	4.094	4.647	5.245	5.797
75.000	2.902	3.219	4.036	4.580	5.169	5.713
80.000	2.861	3.174	3.978	4.515	5.096	5.632
85.000	2.821	3.130	3.923	4.452	5.025	5.554
90.000	2.782	3.0 86	3.869	4.391	4.956	5.477
95.000	2.744	3.044	3.816	4.331	4.888	5.403
100.000	2.708	3.004	3.765	4.273	4.823	5.330
105.000	2.672	2.964	3.715	4.216	4.759	5.260
110.000	2.637	2.925	3.667	4.161	4.697	5.191
115.000	2.603	2.888	3.620	4.108	4.636	5.124
120.000	2.570	2.851	3.574	4.055	4.577	5.059
125.000	2.537	2.815	3.529	4.004	4.520	4.996
130.000	2.506	2.780	3.485	3.955	4.464	4.934
135.000	2.475	2.746	3.442	್. 906	4.409	4.873
140.000	2.445	2.713	3.401	3.8 5 9	4.356	4.814
145.000	2.416	2.680	3.360	3.813	4.304	4.757
150.000	2.387	2.649	3.320	3.768	4.253	4.700
155.000	2.360	2.618	3.281	3.724	4.203	4.645
160.000	2.332	2.587	3.243	3.681	4.155	4.592
165.000	2.306	2 .558	3.206	3.639	4.107	4.539
170.000	2.280	2.529	3.170	3.598	4.061	4.488
175.000	2.254	2.501	3.135	3.558	4.015	4.438
180.000	2.229	2.473	3.100	3.518	3.971	4.389
185.000	2.205	2.446	3.066	3.480	3.928	4.341
190.000	2.181	2.420	3.033	3.442	3.885	4.294
195.000	2.158	2.394	3.001	3.405	3.844	4.248
200.000	2.135	2.369	2.969	3.369	3.803	4.204
205.000	2.113	2.344	2.938	3.334	3.7 6 3	4.160
210.000	2.091	2.320	2.908	3.300	3.724	4.116
215.000	2.069	2.296	2.878	3.266	3.686	4.074
220.000	2.048	2.273	2.849	3.233	3.649	4.033

NWC TP 6525

CON	VERSION FA	ACTORS FOR	R P=	585		TORR
		= X MG/CL		TABLE		
T(C)	C10H8	C11H10	C14H10	C15H10	C19H12	C20H12
20.000	4.098	4.547			7.300	8.069
25.000	4.030	4.470	5.604	4.359	7.178	7.933
30.000	3.9 6 3	4.397		€.254	7.059	7.802
35.000	3.899		5.422	5.153	5.945	7.676
40.000	3.836		5.335		6.834	7.553
45.000	3.776	4.189			5.725	7.434
50.000	3.718	4.124			6.622	7.319
55.000	3.661	4.061	5.091		5.521	7.208
60.000	3.606	4.000	5.015	5.691	6.423	7.099
65. 000	3.553	3.941	4.940	5.507	5.328	5.994
70.000	3.501	3.884	4.868	5.525	٤.23٤	6.892
75.000	3.451	3.928	4.798	5.445	5.146	5. 793
80.000	3.402	3.774	4.731	5.368	6.059	6.697
85.00	3.354	3.721	4.564	5.293	5.975	5.504
90.000		3.670			5.892	6.513
95.000	3.263	3.520	4.538	5.150	5.812	5.424
100.000	3.219	3.571	4.477	5.080	5.734	6.338
105.000	3.177	3.524	4.419	5.013	5.559	6.254
110.000	3.135	3.478	4.360	4.948	5.585	6.173
115.000	3.095	3.433	4.304	4.884	5.513	5.093
120.000	૩.૦55	3.390	4.249	4.822	5.443	6.015
125.000	3.017	3.347	4.196	4.761	5.374	5.940
130.000	2.980	3.306	4.144	4.702	5.308	5.866
135.000	2.943	3.265	4.093	4.645	5.242	5.794
140.000	2.908	3.226	4.043	4.588	5.179	5.724
145.000	2.873	3.187	3.995	4.534	5.117	5.656
150.000	2.839	3.149	3.948	4.480	5.057	5.589
155.000	2.806	3.112	3.902	4.428	4.998	5.524
160.000	2.773	3.077	3.857	4.377	4.940	5.460
165.000	2.742	3.041	3.812	4.327	4.883	5.397
170.000	2.711				4.828	
175.000	2.580	2.974	3.727	4.230	4.774	
180.000	2.451	2.941	3.686	4.183	4.722	5.219
185.000	2.622	2.909		4.138	4.670	
190.000	2.594	2.877	3.607	4.093	4.620	5.106
195.000	2.566	2.846	3.568	4.049	4.570	5.051
200.000	2.539	2.816	3.530	4.006	4.522	4.998
205.000	2.512	2.787	3.493	I.964	4.475	4.946
210.000	2.486	2.758	7.457	7.927	4.428	4.895
215.000	2.461	2.730	J.422	J.98J	4.383	4.844
220.000	2.436	2.702	3.387	I.844	4.339	4.795

NWC TP 6525

CONV	ERSION FA	ACTORS FOR	R 19≔	700		TORR
		= X MG/CL				
T(C)	C10H8	C11H10	C14H10	C16H10	C18H12	C20H12
20.000	4.904	5.440	6.820	7.739	8.735	9.655
25.000	4.822	5.349	6.705	7.609	8.589	9.493
30.000	4.742	5.261	6.595	7.484	8.447	9.336
35.000	4.565	5.175	6.487	7.362	8.310	9.185
40.000	4.591	5.093	6.384	7.245	8.177	9.038
45.000	4.518	5.013	6.283	7.131	8.048	8.896
50.000	4.448	4.935	6.186	7.020	7.924	8.758
55.000	4.381	4.860	6.092	5.913	7.803	8.524
40.000	4.315	4.787	6.000	6.8 09	7.686	8.495
65. 000	4.251	4.716	5.912	6.709	7.572	8.369
70.000	4.189	4.647	5.825	6.611	7.462	8.247
75.000	4.129	4.581	5.742	6.516	7.355	8.129
80.000	4.070	4.516	5.660	6.424	7.250	8.014
85.000	4.014	4.453	5.581	6.334	7.149	7.902
70.000	3.958	4.391	5.505	6.247	7.051	7.793
95.000	3.905	4.332	5.430	6.162	6.955	7.687
100.000	3.852	4.274	5.357	6.079	6.862	7.584
105.000	3.801	4.217	5.286	5.999	5.771	7.484
110.000	3.752	4.162	5.217	5.921	6.683	7.386
115.000	3.703	4.108	5.150	5.844	6.596	7.291
120.000	3.656	4.056	5.084	5.770	6.513	7.198
125.000	3.610	4.005	5.020	5.697	6.431	7.108
130.000	3.545	3.955	4.958	5.627	6.351	7.019
135.000	3.522	3.907	4.897	5.558	6.273	6.933
140.000	3.479	3.860	4.838	5.490	6.197	6.849
145.000	3.437	3.813	4.780	5.425	6.123	5.768
150.000	3.397	3.768	4.724	5.361	6.051	6.688
155.000	3.357	3.724	4.669	5.298	5.980	6.609
160.000	3.318	3.681	4.615	5.237	5.911	6.533
165.000	3.281	3.639	4.562	5.177	5.843	6.459
170.000	3.243	3.598	4.510	5.119	5.777	6.386
175.000	3.207	3.558	4.460	5.061	5.713	6.314
180.000	3.172	3.519	4.411	5.006	5.650	6.24 5
185.000	3.137	3.480	4.363	4.951	5.588	6.176
190.000	3.103	3.443	4.316	4.898	5.528	6.110
195.000	3.103	3.406	4.270	4.845	5.469	6.045
200.000	3.070 3.03 8	3.370	4.270		5.411	
205.000				4.794		5.981
210.000	3.006 2.975	3.33 5 3.300	4.180	4.744	5.3 5 4	5.918
215.000			4.137	4.695	5.299	5.857
	2.944	3.266	4.095	4.647	5.245	5.797
220.000	2.915	3.233	4.053	4.599	5.192	5.738

NWC TP 6525

CON	VERSION FA	ACTORS FOR	R F=	760		TORR
	1PPM	= X MG/CL	J-M FROM			
T(C)	C10H8	C11H10	C14H10	C16H10	C18H12	C20H12
5 0.000						
20.000	5.324	5.907	7.404	8.402	9.484	10.482
25.000	5.235	5.808	7.280	8.261	9.325	10.306
30.000	5.149	5.712	7.160	8.125	9.171	10.136
35.000	5.065	5.619	7.044	7.993	9.022	9.972
40.000	4.984	5.529	6.931	7.866	8.878	9.812
45.000	4.906	5.442	6.822	7.742	8.738	9.458
50.000	4.830	5.358	6.716	7.622	8.603	9.509
55.000	4.756	5.276	6.614	7.506	8.472	9.364
40.000	4.685	5.197	6.515	7.393	8.345	9.223
45.000	4.615	5.120	6.418	7.284	8.221	9.087
70.000	4.548	5.046	6.325	7.178	8.101	8.954
75.000	4.483	4.973	6.234	7.074	7.985	8.826
80.000	4.419	4.903	6.146	6.974	7.872	8.701
85. 000	4.358	4.834	6.060	6.877	7.762	8.579
90.000	4.298	4.768	5.976	6.782	7.655	2.461
95.000	4.239	4.703	5.895	5.690	7.551	8.346
100.000	4.182	4.640	5.816	6.600	7.450	8.234
105.000	4.127	4.578	5.739	6.513	7.351	8.125
110.000	4.073	4.519	5.664	6.428	7.255	8.019
115.000	4.021	4.460	5.591	6.345	7.162	7.916
120.000	3.970	4.404	5.520	6.264	7.071	7.815
125.000	3.920	4.348	5.451	6.186	6.982	7.717
130.000	3.871	4.294	5.383	6.109	6.875	7.621
135.000	3.824	4.242	5.317	6. 034	5.811	7.528
140.000	3.777	4.190	5.253	5.961	6.728	7.437
145.000	3.732	4.140	5.190	5.890	5.648	7.348
150.000	3.688	4.091	5.129	5.820	6.569	7.261
155.000	3.645	4.044	5.069	5.752	6.493	7.176
160.000	3.603	3.997	5.010	5.686	6.418	7.093
165.000	3.562	3.951	4.953	5.621	6.344	7.012
170.000	3.521	3.907	4.897	5.557	6.273	6.933
175.000	3.482	3.863	4.842	5.495	6.203	6.856
180.000	3.444	3.820	4.789	5.435	6.134	6.780
185.000	3.406	3.779	4.737	5.375	6.067	6.706
190.000	3.369	3.738	4.686	5.317	6.002	6.633
195.000	3.333	3.698	4.635	5.260	5.938	გ.563
200.000	3.298	3.659	4.586	5.205	5.875	6.493
205.000	3.264	3.621	4.539	5.150	5.813	5.425
210.000	3.230	3 .58 3	4.492	5.097	5.753	6.359
215.000	3.197	3.546	4.446	5.045	5.694	6.294
220.000	3.164	3.510	4.400	4.994	5.636	6.230

Appendix E

SPEX FLUORIMETER CORRESPONDENCE



April 19, 1983

Dr. James Short
Rural Route #1
Box 197
Switz City, Indiana 47465

Dr. Richard Loda Naval Weapons Center Mail Code 3851 China Lake, CA 93555

Dear Drs. Short and Loda:

I am sending each of you a copy of this letter and duplicate information packet concerning our FLUOROLOG 2 spectrofluorometers as the same instrument applies to each of you.

When looking at PAH gasses from burning flares, extreme sensitivity will be required due to the low molecular densities involved. You will note that with fluorescein 0.1 ppt is detectable even with our least expensive model 111C instrument with single monochromators, room temperature PMT and 150 watt Xe lamp. Even better minimum levels of detectability are possible with our model 112 instrument which might include a cooled PMT and 450 watt lamp. Single photon counting detection, efficient classically ruled gratings in our monochromators, and very good 1:1 imaging optics account for much of the sensitivity of the FLUOROLOG 2 instruments. By taking these values and backtracking to the vapor pressures included in the burning flares, your 1 ppm levels should be easy to achieve.

The Datamate scan controller, photometer, and data system included with the FLUOROLOG 2 instruments is quite easy to use despite its initial apparent complexity. Being microprocessed, default values are loaded into the Datamate on start up making the taking of data easy. As required, additional steps can be added to the data acquisition such as repetitive scanning, peak location, excitation and emission correction, etc. Once the appropriate steps are determined, a simple program can be developed by simply telling the Datamate to perform the various steps in sequence. This is called keystroke programming. You will note in some of the attached Tech Notes on the Datamate how it is not only easy to use but can be quite powerful.

Drs. Short and Loda April 19, 1983

Delivery and installation of an instrument will probably not be a limiting problem in your possible purchase of an instrument. This is assuming that Spex will be informed as the purchase of an instrument progresses. The mentioned mid-June delivery and installation at Los Alamos is well within our capabilities.

I hope this information helps in your determination of the suitability of our FLUOROLOG 2 instruments in your work. I will contact Richard and make arrangements for samples to be run either in Los Angeles, Santa Barbara, the San Francisco Bay Area, or in our New Jersey factory.

In the meantime, if either of you have any questions concerning the FLUOROLOG 2, feel free to give me a call.

Sincerely,

SPEX INDUSTRIES, INC.

Hollis F. Davis

Western Regional Manager

ps

Attachments



29 April 1983

Dr. Richard Loda Naval Weapons Center Code 3851 China Lake, CA 93555

Dear Dr. Loda:

Enclosed are the fluorescence spectra of gas phase phenanthrene and pyrene which you requested. The spectra were obtained on a FLUOROLOG F112 system with a 450W Xenon lamp, and a cooled R928 detector. The price of this system with a dual disk drive is approximately \$36000. Hollis Davis can discuss prices with you in greater detail.

The operating parameters are as follows: for pyrene excitation spectrum, emission wavelength = 400 nm, 5 and 10 nm bandpass for excitation and emission monochromators respectively, a 1 nm step size and 1 second integration time. For pyrene emission spectrum 321 nm excitation wavelength, 10 and 5 nm bandpass, 1 nm step size and 1 second integration time. For phenanthrene excitation spectrum, 366 nm emission wavelength, 5 and 10 nm bandpass, 1 nm step size and 1 second integration time. For phenanthrene emission spectrum, 241 nm excitation wavelength 10 and 5 nm bandpass, 1 nm step size, and 1 second integration time.

Plot A shows uncorrected (dashed trace) and corrected excitation spectra of pyrene at 60°C. The correction was done by a rhodamine B quantum counter. Plot B is uncorrected (solid line) and corrected emission spectra of pyrene at 60°C. The instrument response function for corrected emission spectra was determined with a standard lamp. Plots C and D are excitation and emission spectra at 60°C of phenanthrene.

Because of the signal levels, I did not think that synchronous scanning would be very informative. Therefore, I obtained emission spectra of a mixture of phenanthrene and pyrene exciting at two different excitation wavelengths (241 and 321 nm). I performed the above experiment at 60, 50 and 30°C. The results are given in Plots E-J.

Dr. Richard Loda 29 April 1983 Page 2

I think you will agree that these results clearly show that our instrument is capable of detecting PAH's in the gaseous state at the part per million level. If you have any questions about the data, please contact me.

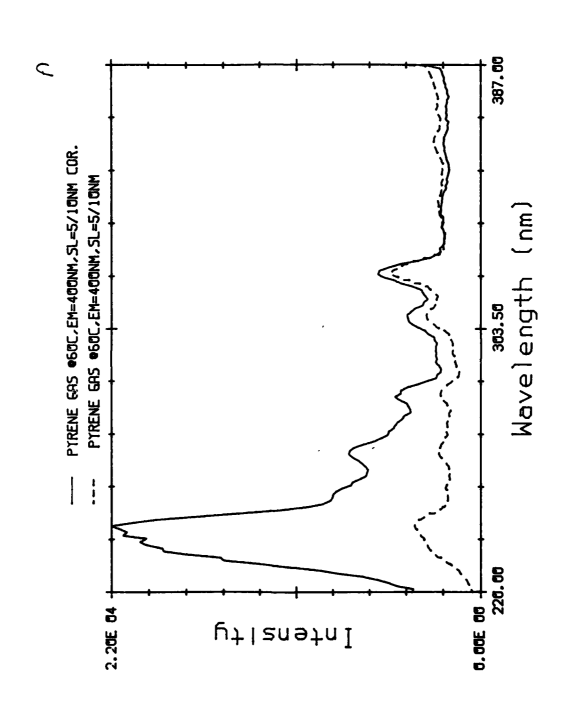
Sincerely yours,

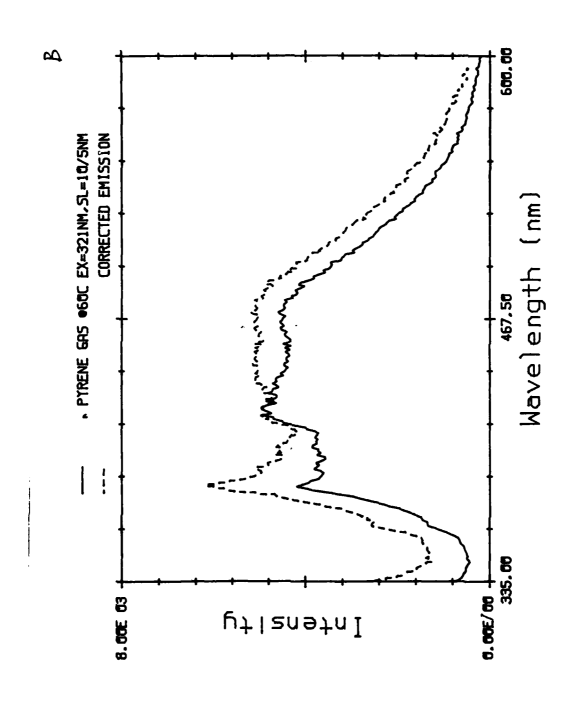
SPEX INDUSTRIES, INC.

Frank Purcell, Ph.D. Applications Manager

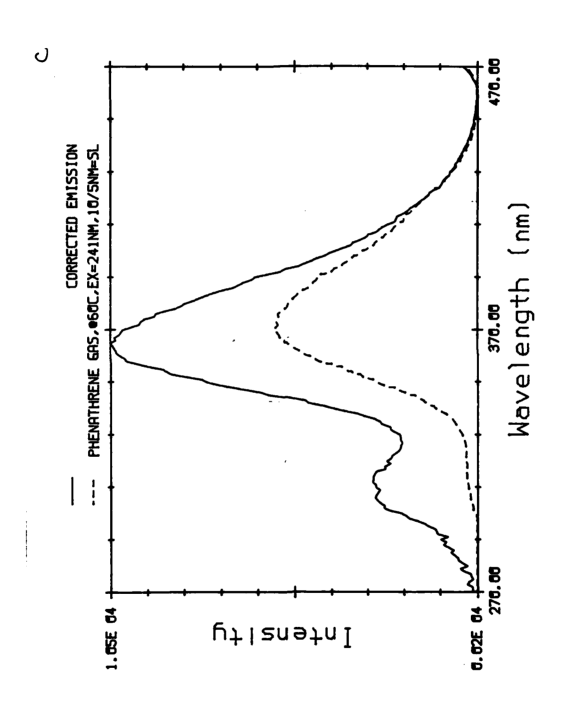
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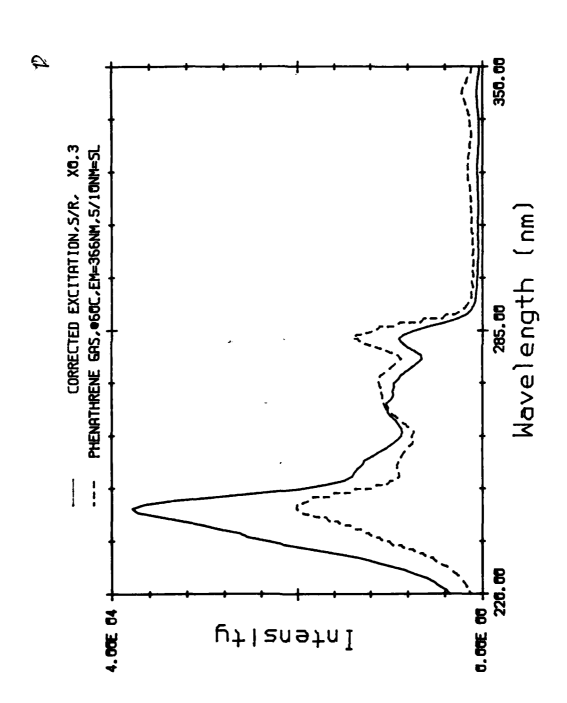
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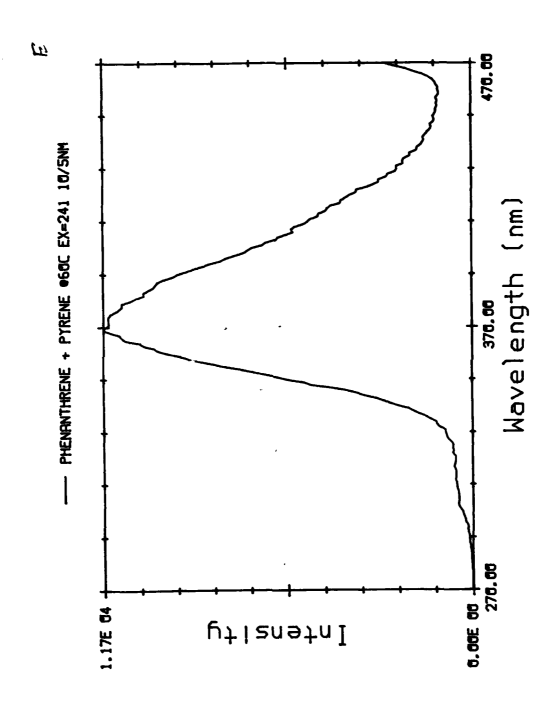


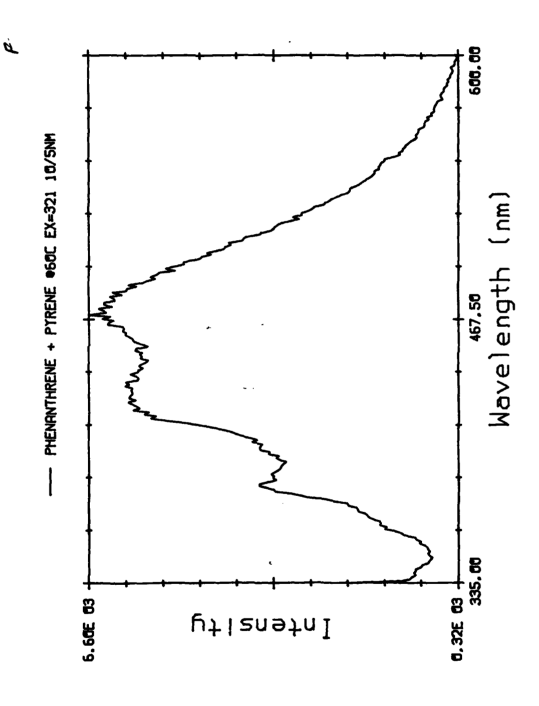


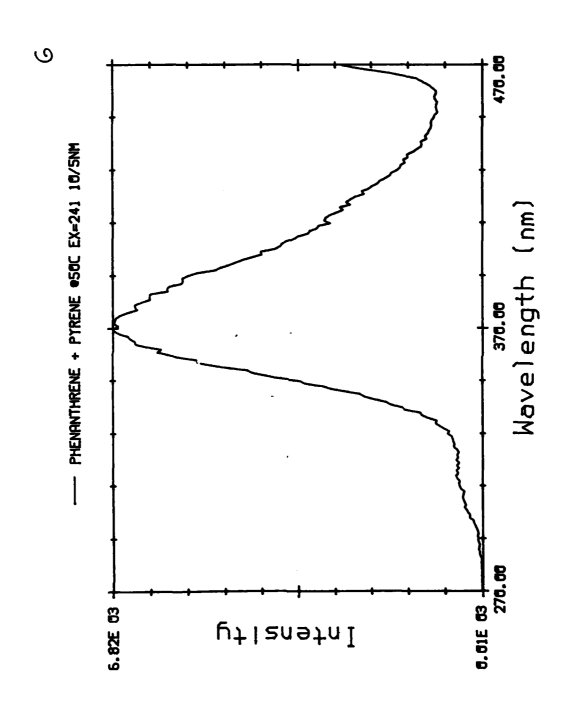
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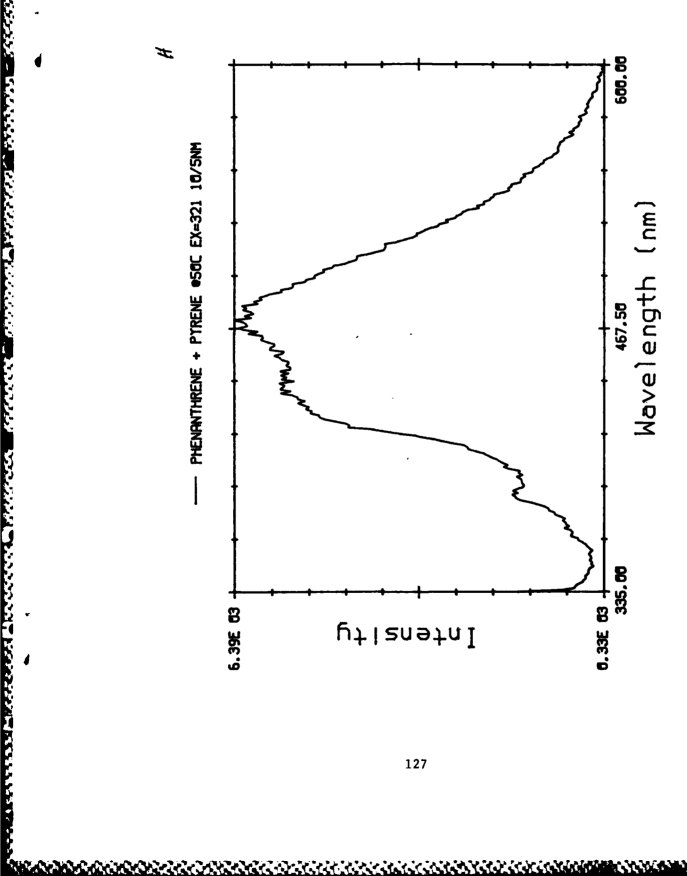


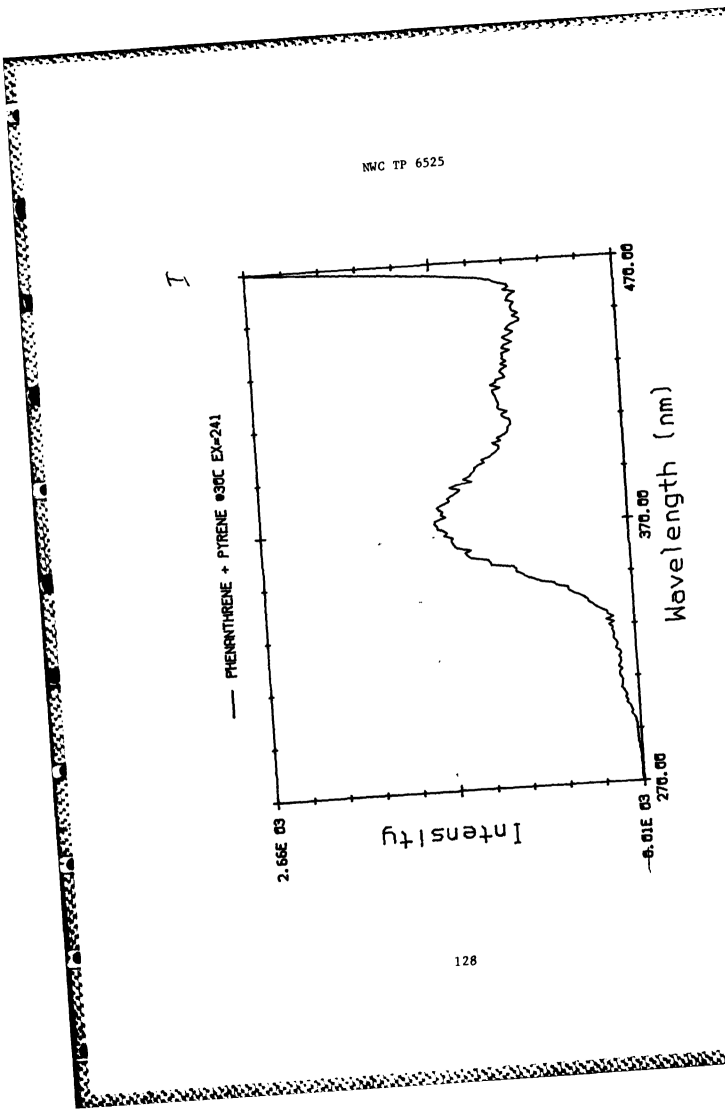


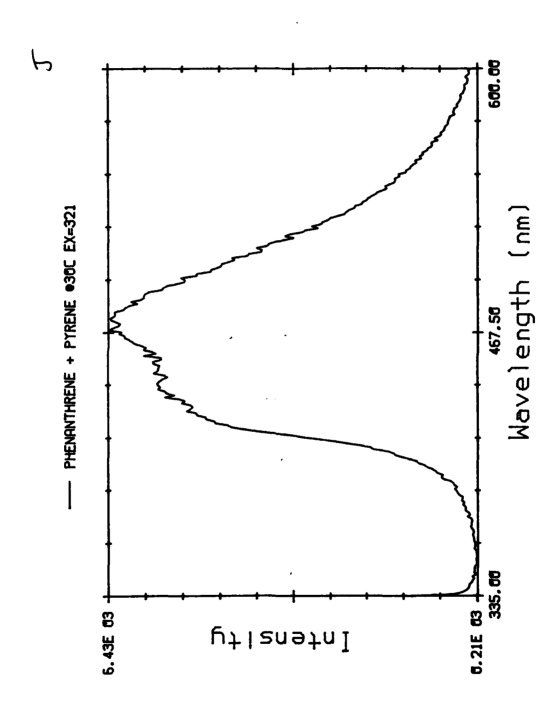


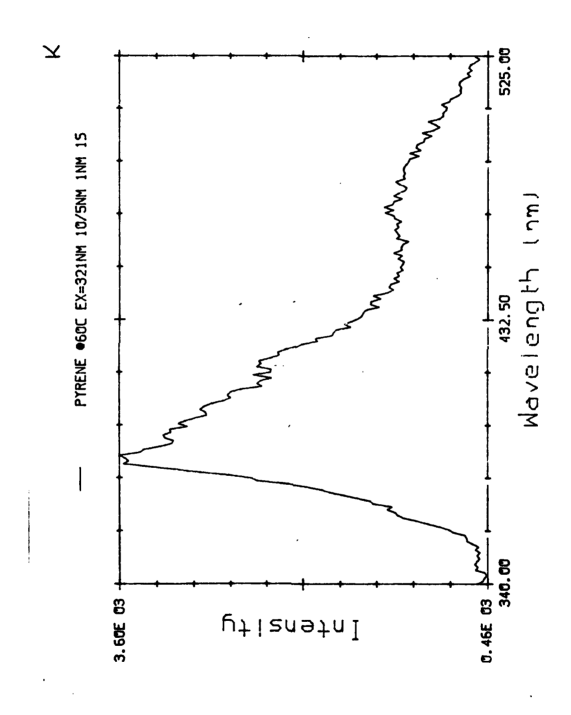


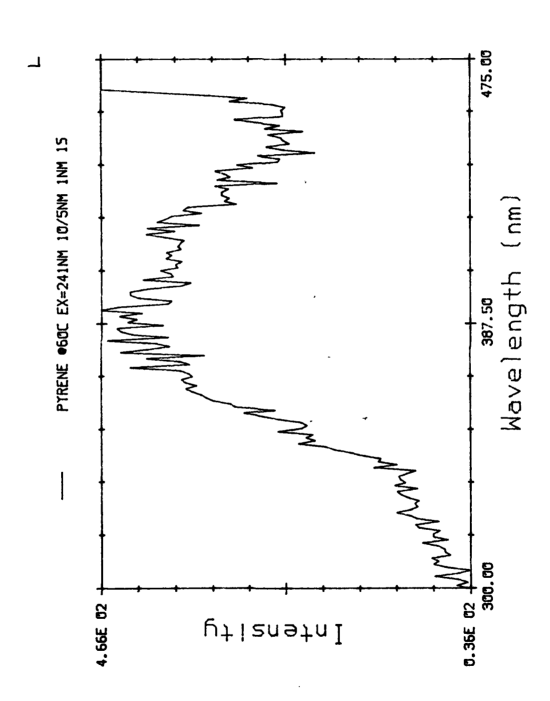
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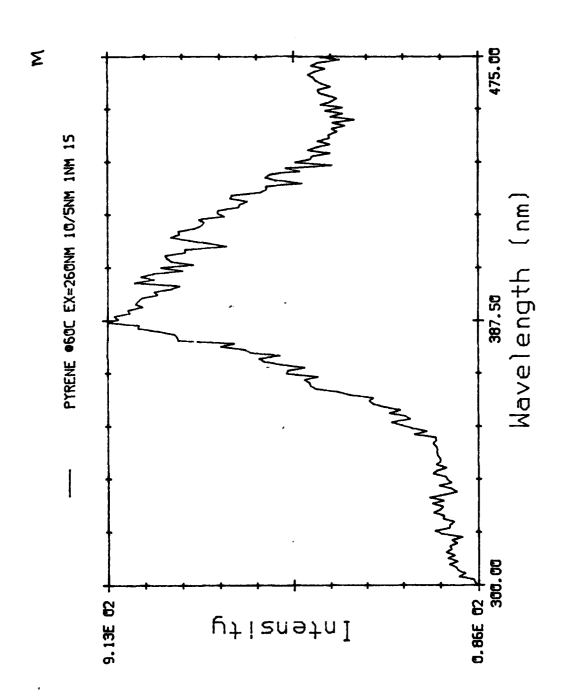


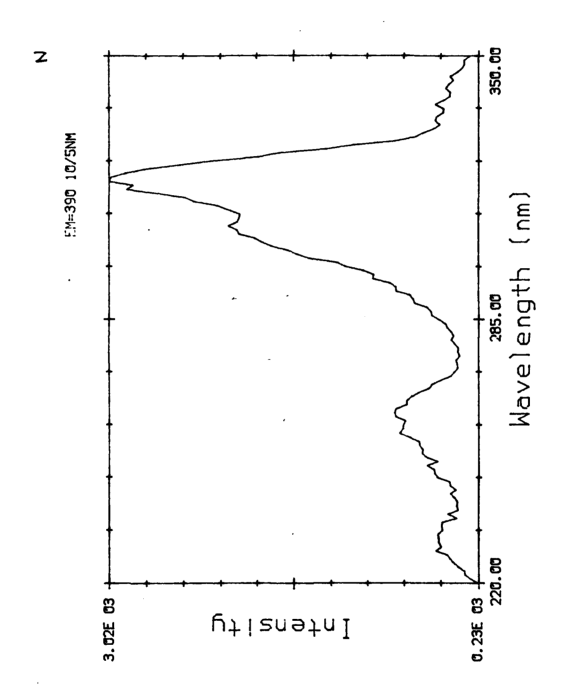












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